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MEASURING FOOD VALUES FOR DAIRY COWS1,2

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THE EVALUATION OF FEEDS

FEED ACCOUNTS for over half the total cost involved in animal production (Hopkins, 1940). Obviously, then, it is important to evaluate feeds correctly.

Replacement Equivalent.—The problem can be best introduced by a simple example: If a dairy farmer can secure corn for \$38 a ton or wheat for \$37 a ton, which is the better buy?

To answer this question the farmer must know how much wheat will replace in his ration a ton of corn so that the result—milk yield and change of body substance in the cows—will remain the same.

In this paper the figure that indicates how much of one feed is necessary to replace a certain amount of another feed is called *replacement equivalent*. Since, as a rule, a ration consists of several feeds, the farmer is especially interested in knowing the replacement equivalent of feeds when they form only a part of a ration. If, for example, a daily ration includes 10 pounds of hay, 1 pound of beet pulp, 1 pound of cottonseed meal, and 3 pounds of corn, how much wheat would be required to replace these 3 pounds of corn so that the new ration with wheat has the same food effect as the old ration with corn? In this case, as usually in food evaluation, only a part of the ration is replaced; and we therefore speak here of *partial replacement equivalents*.

Data Available at Present.—Replacement equivalents can be derived from the standard tables on total digestible nutrients: digestible protein plus digestible carbohydrates plus (2.25 × digestible fat). As Morrison (1936) shows, corn contains 83.7 pounds of total digestible nutrients per 100 pounds, and wheat 83.6. This is 94.6 pounds of total digestible nutrients per 100 pounds of dry matter for corn, 93.8 for wheat. Assuming that the offered corn and wheat, in the example cited in the preceding paragraph, have the same moisture content, the wheat is a slightly better buy: it contains 99 per cent of the total

digestible nutrients of corn, and it costs only $\frac{37}{38} \times 100 = 97$ per cent of the price of corn.

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^{*}See "Literature Cited" at end of the paper for complete data on citations, referred to in the text by author and date of publication.

One may calculate the replacement equivalent of corn and wheat also on the basis of their net-energy content and derive from Armsby's (1922) tables a net-energy content of 95.5 "therms" per 100 pounds of dry corn and of 102.2 "therms" per 100 pounds of dry wheat. Here 100 pounds of wheat replaces 107 of corn, so that wheat at a lower price than corn is a considerably better buy. The net-energy value given in an earlier publication by Armsby and Fries (1916), based on the same data for the composition of corn (dent), is almost 5 per cent higher than that mentioned above. The discrepancy probably results from a recalculation on the basis of some different "correction." (See footnote 7, p. 513.)

A third method for calculating replacement equivalents of feeds may be based upon Kellner's (1919) tables of *starch values*. In the "production feed" for fattening steers, 100 kg of dry corn replaces 93.8 kg of starch; 100 kg of dry wheat replaces 80.5 kg of starch. Consequently 100 kg of dry wheat re-

places $\frac{80.5}{93.8} \times 100 = 86$ kg of dry corn. From this calculation the farmer con-

cludes that corn at \$38 per ton is a considerably better buy than wheat at \$37 per ton—just the opposite of the conclusion based on Armsby's net energy!

Further, according to Möllgaard's (1929) table on dairy feeding, 1 kg corn represents 2.17 Möllgaard feed units, and 1 kg wheat represents 2.00. (One Möllgaard feed unit equals 1,000 kilocalories net energy for milk production.) On the basis of these figures, 100 kg wheat replaces 92 kg corn, and corn is a better buy. The Möllgaard feed units cannot, however, be regarded as an independent type of feed evaluation: Möllgaard based his feed units, not on his own respiration trials with dairy cows, but on Kellner's respiration trials with fattening steers. The discrepancy between the replacement equivalent calculated from the Möllgaard feed units and that calculated above from Kellner's starch values therefore probably resembles the discrepancy between Armsby's net energy for corn in 1922 and that in 1916, because it probably results from different "corrections" applied to the same original data.

The Scandinavian feed units offer a fourth possible way of calculating the nutritive replacement equivalent of wheat for corn. These are replacement equivalents with 100 kg barley as the standard unit of reference. Thus in Nils Hansson's (1928) table, one finds that in dairy rations 100 kg corn can replace 105.7 kg barley, and 100 kg wheat can replace 101.1 kg barley. So 100 kg wheat

replaces $\frac{101.1}{105.7} \times 100 = 95.6$ kg of corn. This comparison makes corn at \$38

a ton a slightly better buy than wheat at \$37 a ton.

According to total digestible nutrients and Armsby's net energy, wheat is a better buy; but according to starch value and Scandinavian feed units, corn is the cheaper feed. This contradictory conclusion is based on the replacement equivalent of wheat for corn, summarized as follows for the four systems of food evaluation: according to total digestible nutrients, 100 kg wheat replaces 99 kg corn; with Armsby's net energy, 100 kg wheat replaces 107 kg corn; with Kellner's starch value, 100 kg wheat replaces 86 kg corn; and with Scandinavian feed units, 100 kg wheat replaces 96 kg corn.

Critique of Total Digestible Nutrients, Net Energy, and Starch Value as Bases for Evaluating Dairy Feeds.—To inquire which of these four answers is correct would be uncritical; but one may reasonably ask which is most nearly correct for the problem under discussion.

To calculate replacement equivalents on the basis of total digestible nutrients involves the assumption that 1 calorie of digestible energy in a food has the same nutritive value as 1 calorie of digestible energy in any other food—an assumption that has been challenged, at least in comparisons between roughage and concentrates. Wolff and his co-workers (1888) noticed that the digestible food energy in oats was more efficiently utilized for work of horses than the digestible food energy in hay. Schneidewind and his collaborators (1910) in their paired trials on steers also indicated that digestible food energy in corn has a greater fattening effect than digestible energy in hay.

Net energy expresses the energy which appears in the animal's products or which serves to keep the animal alive. It is the only unquestionably correct measure for the useful food energy in any one case for which it has been measured, at least if the result has not been subjected to so-called "corrections" in an attempt to make them applicable to conditions other than those of the measurements. How, then, could Meigs (1925) conclude that "net energy values are a less accurate measure of the relative values of feeds for maintaining uniform body weight than are total digestible nutrients"?

In his critique of net energy Meigs questioned, not the correctness of the net energy as measured directly in any one trial, but the generalization from a few results. As he showed, even with the same food, clover hay, for example, the net energy amounted to 51 per cent of the metabolizable energy in one trial, to 78 per cent in another, and to 92.7 per cent in a third trial. With such variability in the results for one single feed, measured with all possible care under well-controlled conditions, how can one expect to derive from a few trials a satisfactory mean, and even to calculate from it net-energy values for those feeds that have not been tested? Considering this great variability of the bases of Armsby's calculation, one can easily understand that in some trials, such as that of Meigs, metabolizable energy may happen to fit the results better (that is, express the true net energy of the feed under the particular conditions better) than does the net energy derived from Armsby's calculation (1922, p. 674). One is, of course, not justified in concluding, from any such single case, that Armsby's calculated net-energy values are in general less reliable measures for food values than is metabolizable energy, especially

Tealeulating, for example, an animal's heat production to a standard day of 12 hours standing and 12 hours lying is an attempt to apply the results to conditions different from those under which they were obtained. Even if the energy cost of standing over lying were known satisfactorily, this calculation might be the opposite of what is generally understood by the term "correction." Some feeds, as Zuntz and his co-workers (1931, p. 784) report, may increase the "Bewegungstrieb" of animals and thus waste energy; others, such as alcohol, may induce the animal to lie down longer than with other feeds and thus save energy. To discover that the utilization of food energy is related to such effects of feeds on the nervous system would be of great interest; but to calculate the heat production of the animals to a standard day of 12 hours standing would eliminate rather than bring out such particular effects on food utilization. Certainly the claim would be unjustified that theoretical feed values obtained by eliminating mathematically the particular energetic effects of feeds are more correct than the feed values directly measured under the actual conditions of the experiment.

when the latter is also merely calculated. The variability in the composition of feeds, even between feeds that are assumed to be the same, is presumably a major source of discrepancy in results. (See discussion of the tabulation

on p. 520.)

Mitchell's explains the great variability in Armsby's results by the particular experimental technique and evaluation of results followed in Armsby's earlier trials, including an exaggerated correction for standing over lying. The more recent animal-calorimeter measurements at Pennsylvania State College show a smaller variability (compare, however, footnote 22, p. 550). Mitchell (1934) himself, however, advanced the hypothesis that "except for differences in digestibility the net energy value of all perfectly balanced rations is the same under the same conditions of feeding."

Some—at least theoretical—difficulty with the strict proof of Mitchell's hypothesis is the introduction of a none-too-well-defined limiting condition: "perfectly balanced ration." If the rations to be compared need be balanced only with regard to the protein-to-energy ratio, then the problem can be readily attacked as Schneidewind and his co-workers (1910) attacked it in their experiments with steers. If, however, following Axelsson's (1939) idea, the rations must also be equally balanced with respect to the fiber content, the task becomes already rather complicated. And if we go a step further, then it becomes impossible to disprove Mitchell's hypothesis, because any deviation from the expected result could be explained by an imbalance of the rations with regard to some other, perhaps not yet known, nutritional factor.

Considering the problematic nature of a well-balanced ration, we are tempted to reverse the formulation of the hypothesis to read: "For a given production two rations are equally well balanced when the partial efficiency (or net availability, in Mitchell's terms) of the metabolizable energy for that production is equal." Instead of making a measurable quantity (efficiency) dependent on an unknown (balance of ration), this formulation defines the

unknown (balance) in terms of the measurable (efficiency).

Mitchell's hypothesis would mean that differences in calorigenic action or in the partial efficiency (or net availability) of metabolizable energy are only a matter of balance of the ration. If this is generally true, then indeed feeds can be evaluated according to their content of metabolizable energy, just as Meigs (1925) proposed—a proposal apparently accepted by Kriss and Forbes in their new method of deriving a feeding standard for dairy cows (Kriss, 1931), which is a return to the system of evaluation by total digestible nutrients.

Kellner's trials (1896, 1900) are somewhat more suitable than Armsby's for deriving net-energy values of untested feeds: as reference substances he used semipure nutrients, especially starch, gluten, and oil, so that he could base the calculation for untested feeds on their content of digestible protein, fat, and carbohydrates, whereas Armsby disregarded the chemical composition. Much of Meigs' criticism, however, as he himself states, applies to Kellner's results as well as to Armsby's; and a later review by Axelsson (1939) reveals many defects in some of Kellner's basic work.

The comparison (made on page 512) of the replacement equivalents of wheat

⁸ Personal communication of H. H. Mitchell with the senior author, 1943.

for corn according to Armsby (107 kg) and according to Kellner (86 kg), both based on similar experiments with steers, is not encouraging for applying either of these systems directly in evaluating feeds for dairy cows. If this application is made, as by Möllgaard (1929), then upon the uncertainty in the list of starch values for fattening is superimposed a dubious assumption—namely, that the relative food values are the same for functions physiologically so different as fat deposition and lactation.

Evaluation of feeds for dairy cows on the basis of Scandinavian feed units has one great advantage over the total digestible nutrients, net energy, and starch value: the Scandinavian feed units are based on trials with dairy cows. Such is not the case for those figures which were based on Hansson's (1914) "milk-starch values" assuming that 1 feed unit equals 0.75 milk-starch value. The milk-starch values of feeds are calculated from their content of digestible nutrients as in the calculation of Kellner's starch values; but for 1 kg digestible protein an equivalent of 1.43 kg starch is used instead of Kellner's 0.94, which latter equivalent is correct only when protein is used for fat production. (Hansson's protein factor has been discussed by Kleiber, 1929.) When, however, the result of a group trial differed from the calculated "milk-starch values"—for example, from the value for wheat bran—Hansson relied on the result of the group trial.

Assuming that the feed units for wheat and corn used above are the results of Scandinavian group trials, we consider the equivalent of 96 kg corn for 100 kg wheat to be the most reliable of the four figures as a basis for estimating the relative food value for dairy cows. Yet the Scandinavian feed units themselves are open to criticism.

Critique of the Scandinavian Feed Units.—Some of the criticism against the Scandinavian feed units is based on a misunderstanding of the meaning of replacement equivalent, as shown earlier by Kleiber (1929). Two major criticisms, however, are justified:

- 1. In the Scandinavian group trial, body weight serves as index for body substance. The trials, especially in recent times, are so conducted that only minor changes in body weight occur. A cow can, however, maintain her body weight and yet lose body fat, because she can replace the weight of fat by the weight of water. The effect of this uncertainty on the result is not likely to be 100 per cent, as Möllgaard (1927) suggested; but it may cause considerable error.
- 2. Even though the exchange of barley with wheat or corn offers no essential difficulty, the replacement of barley as the reference substance by a feed rich in protein (such as cottonseed meal) leads to a change in the protein:energy ratio in the ration. Such a change in the protein:energy ratio may, however, affect the efficiency of energy utilization, as Möllgaard has shown (1923, 1929); his results on dairy cows have been confirmed on rats by Mitchell (1934), by Hamilton (1935), and by Forbes with his co-workers (1935). This problem has been reviewed by Mitchell (1937). Mitchell, Hamilton, and Haines (1940) found, however, no effect of inadequate protein supply (imbalance of ration) on the energy utilization of young steers. In practical group trials one may possibly keep the protein:energy ratio in a barley ration and in the corresponding ration with cottonseed meal within a range in which

changes of this ratio have but an insignificant effect on the efficiency of energy utilization. According to Möllgaard (1931, p. 327) this range extends from a production quotient of 0.14 to 0.23. (Möllgaard's production quotient is the

ratio of protein net energy to total net energy for fattening.)

Even here, however, it is unsatisfactory to compare feeds having different protein content with one given reference substance, such as barley or starch. Protein and energy have entirely different functions in the animal body; they can therefore not be reduced to a single physiological basis of comparison. This fact has been recognized by many workers in this field; Petersen (1932) correctly calculates feed values from digestible energy and protein, using as a basis the prices of two standard feeds, namely cottonseed meal and corn, instead of one.

Purpose of the Present Trials.—In the experiments reported here, the aim was to investigate the possibilities of improving the evaluation of feeds. We were not encouraged by Forbes's belief (1932) that "there is no present, or possible [!] scientifically significant and defensible single measure of nutritive value of individual feeding stuffs." Obviously, however, Forbes is thinking in terms of absolute, not relative, feed values; and he does not consider the concept of partial effects, a concept very useful even when such partial effects cannot be measured directly. One can derive partial molal volumes of solutes, for example, only by measuring volumes of entire solutions; yet physical chemists do not conclude that partial molal volumes are "insignificant" or "indefensible." Likewise, one may advantageously use partial nutritive effects even though only total food effects are (at present) directly measurable.

If a complete ration of 10 pounds of hav and 3 pounds of wheat has the same net energy as another complete ration with 10 pounds of the same hay and 2.9 pounds of corn, then the partial nutritive effect (with respect to energy) of 3.0 pounds of wheat is equal to that of 2.9 pounds of corn; or the partial replacement equivalent of 1 pound of wheat is 0.96 pound of corn. Our trials were started in the belief that to measure such partial replacement equivalents is practically possible and scientifically sound. We realize, of course, that in the measurement of partial effects in nutrition the ideal of changing only one independent variable (such as the amount or the composition of the food) to measure the corresponding partial change in the dependent variable (such as yield of nitrogen or of energy in milk) cannot be accomplished in a strict sense. When we compare the effect of a production ration fed to a lactating cow with that of a maintenance ration fed to a dry cow, not only the food has changed, but also the hormonal condition of the cow. Such hormonal changes are presumably involved in any changes of lactation rates. This unavoidable deviation of the results from ideal partial effects in a strict mathematical sense can, however, hardly lead to confusion or misinterpretation. In the comparison of corresponding partial effects of different feeds as partial replacement equivalents, the influence of these other variables on the results presumably cancels out to a considerable extent. This is one of the major advantages of using a reference substance for the measurements. (Compare p. 549.)

To meet Möllgaard's (1927) major objection against Scandinavian group trials, we measured in a respiration chamber the changes in fat and protein

of the cow's body substance, and thus eliminated the errors involved in estimating these changes from changes in body weight. Judging from our measurements, however, the weight errors involved in a well-conducted Scandinavian group trial may be easily compensated by the larger number of cows, the longer duration of the trial, and the more "normal" conditions as compared with a pair trial in a respiration chamber.

TABLE 1
TIMETABLE OF THE MAIN RESPIRATION TRIALS

Trial no.	Trial period	Level of food intake	Cow no.
		First Series	1
3	Aug. 7 to Aug. 19, 1939	Maintenance on Sudan hav	494, 1003
5	Sept. 4 to Sept. 16, 1939	Maintenance on Sudan hay	494, 1003
7	Oct. 2 to Oct. 14, 1939	Maintenance on Sudan hay	1003, 1009
9	Oct. 30 to Nov. 11, 1939	Maintenance on Sudan hay	1003, 1009
12	Dec. 4 to Dec. 22, 1939	Lactation on Sudan hay and reference substance	1007, 1021
4.4	T 90 1 F.1 10 1010	reference substance	1007
14	Jan. 29 to Feb. 10, 1940	Lactation on Sudan hay and barley	1021
		reference substance	1007
16	Feb. 19 to Mar. 2, 1940	Lactation on Sudan hay and barley with casein supple-	
9		ment	1021
		reference substance	1021
18	Mar. 25 to April 6, 1940	Lactation on Sudan hay and barley with casein supple-	
		ment	1007
20	April 22 to May 5, 1940	Lactation on Sudan hay and reference substance	1007, 1021
22	May 13 to May 25, 1940	Maintenance on Sudan hay	1007, 1021
23	May 27 to June 1, 1940	Fasting	1007, 1021
		Second series	
24	Jan. 27 to Feb. 15, 1941		728
24	Jan. 27 to Feb. 15, 1941	Maintenance on Sudan hay	732
25	Feb. 24 to Mar. 15, 1941	Half maintenance on Sudan hay	732
20	reb. 24 to mar. 15, 1541	Maintenance on Sudan hay	728
26	Mar. 24 to April 5, 1941	Maintenance on Sudan hay and reference substance	728, 732
27	April 7 to April 19, 1941	Maintenance on Sudan hay and reference substance	728, 732
		reference substance	732
28	June 16 to July 3, 1941	Lactation on Sudan hay and barley with casein supple-	
		ment	728
		reference substance	728
29	July 14 to Aug. 2, 1941	Lactation on Sudan hay and barley with casein supple-	Man
		ment	732

More serious than the weight error in the Scandinavian group trials and their results, the feed units, is the classification of feeds with different protein content on the basis of one single reference substance—for example, barley. Such classification may lead to different protein:energy ratios in test ration and reference ration; and it neglects the fact that energy effect and protein effect are essentially and qualitatively different physiological functions. To overcome this difficulty we used as reference substance a variable mixture of two well-defined nutrients—namely glucose (as representative of "nitrogenfree" food energy) and casein (as representative of protein). Such a reference mixture can be adapted to any food to be tested so that the protein:energy

ratio is exactly the same for test ration and reference ration. The reference substance for corn, for example, was composed of 10 per cent casein and 90 per cent glucose, while the reference substance for cottonseed meal consisted of 51 per cent casein and 49 per cent glucose. The simultaneous measurement of protein and energy utilization leads to a twofold result—namely the "nitrogen-free" glucose equivalent and the casein equivalent of the feed tested. This twofold result properly expresses the two essentially different physiological functions of energy and protein.

So far only trials with barley as the test feed have been carried out on dairy cows. These trials lead one to conclude that 100 grams of dry barley as part of a balanced dairy ration replaced on the average 9 grams of casein plus 100

grams of glucose.

Presumably, this casein-glucose equivalent of barley can be applied over a wider range of conditions than the absolute figures obtained in these trials—namely, the net energy in glucose and barley and the net protein in casein and barley for lactation, as well as the net energy and net protein in Sudan hay for maintaining dairy cows.

Originally it was planned to measure the casein-glucose equivalent of cotton-seed meal in addition to that of barley and then to suggest using a mixture of barley and cottonseed meal as an auxiliary reference substance for group trials on feed values with dairy cows. Since the lack of appetite of the cows was a major difficulty in the trials, one may expect that the more palatable auxiliary reference substance would have decided advantages over the mixture of casein and glucose even though the latter is, of course, chemically better defined.

METHODS USED

Table 1 shows the plan and timetable of the experiment. To establish the maintenance requirement, five main trials were conducted, in the first series, with Sudan hay alone while the cows were dry. Two trials (12 and 20) were carried out with both cows on the reference ration; and in three trials one cow was on the test ration while her pair-mate was on the reference ration. This series was concluded with a fasting trial of 5 days' duration. The effect of Sudan hay between the half maintenance and the maintenance level was studied in the second series. In two trials dedicated to measuring the partial effect of casein and glucose for maintenance, one cow received hay and concentrate together twice a day; the other cow, casein and glucose alone in the morning and hay alone in the evening. The last two trials were run as a combined pair and period trial to compare barley and the corresponding mixture of glucose and casein.

EXPERIMENTAL COWS

A year before the trials began, we experimented with twelve cows in order to study the palatability of various feed combinations and then to train the cows in the intake of the somewhat artificial reference diet. From these twelve animals were selected the pairs for the respiration trial. Some of their characteristics are summarized in table 2. Their mean body weights appear in table 4.

⁹ Net nitrogen (a term analogous to net energy) is that part of the nitrogen in food which either appears as nitrogen of body substance or milk or eggs, or is equal to the loss of body nitrogen prevented by the food. The term "net protein" was first used by Mitchell (1922).

FEED AND RATIONS

Palatability.—A compromise had to be reached between two aims—first, to make the ration as simple, well defined, and reproducible as possible; second, to make it reasonably palatable. For reproducibility, the ideal ration would have consisted of pure chemical substances. Obviously, however, such a ration is useless if the cows do not eat it or if they eat so little that they cannot be considered in a reasonably normal state of nutrition.

The food was composed of a basic part, kept constant throughout the trial, and a variable or replaceable part. The constant part was expected to cover

Cow no.	Breed	Detection	Milk record for 305-day period, at age of two years		Body weight range	
		Date of birth	Milk, ex- pressed in 1,000 kg	Average milk fat, per cent	during the trials (kilograms)	
494	Jersey	May 28, 1930	4.1	6.6	450-500	
1003	Holstein	Oct. 6, 1934	5.4	3.6	480-530	
1009	Holstein	May 6, 1935	4.8	3.8	540-560	
1007	Holstein	Dec. 13, 1934	5.2	3.7	460-480 (430 at end of fast)	
1021	Holstein	July 22, 1936	3.7	3.7	470-500 (440 at end of fast)	
728	Jersey	July 4, 1937	2.8	5.8	350-380	
732	Jersey	Aug. 9, 1937	3.4	5.6	370-430	

only the maintenance requirement; then the entire "production part," which was added to the maintenance food of the cow during lactation, could be replaced and thus used for measuring replacement equivalents.

Constant Part of the Ration.—The constant part of the ration should contain only enough protein to cover the maintenance requirement, so that all the protein required for lactation could be given in the variable part. In addition, the constant part should provide enough bulk to make the rations "normal" and palatable.

A first attempt was made to use a mixture of oat hay and molasses as the constant part of the ration; and a respiration trial was started with two cows on such a mixture as maintenance food after several weeks of training. In the respiration chambers, however, the cows began leaving more and more of this ration in the mangers. Even though the environmental conditions were kept as agreeable as possible and even though there was some feeling of herd security because the two experimental cows could see each other through a window, the confinement seemed to reduce their appetites; apparently a reliable maintenance trial could not be run in the respiration chamber on a feed that did not particularly appeal to them anyway. In seeking a more palatable feed for the basic part of the ration, Sudan hay was chosen because of its relatively low protein content; cows could be trained to eat a considerable amount of the reference food as an addition to this hay.

The Sudan grass was cut before bloom, dried in the sun, and well cured.

The hay was then chopped and blown to a mow, from which it was taken as needed for the feeding trials.

The following tabulation gives the composition of a sample of Sudan hay:

Per cent	Per cent
Water 10.2	Ash total
Crude protein	Ash insoluble in acid 3.9
Ether extract 1.2	Carotene 0.001
Crude fiber	Calcium 0.354
Lignin 13.4	Phosphorus 0.311
Colluloso 25 2	

Originally it was planned to prepare pellets of ground hay as roughage for the entire trial, to insure uniformity of the constant part of the ration. This plan, which would have eliminated considerable error, unfortunately had to be abandoned because of the cost.

Table 3 shows the nitrogen, carbon, and energy content of the hay. The coefficient of variation of the nitrogen content is \pm 9 per cent for eleven samples used in the preliminary trials, and \pm 5 per cent for ten samples of hay used in the main trials. Fortunately the variability of the carbon content was smaller—namely, \pm 1.9 and \pm 1.6 per cent for the samples in preliminary and main trials respectively. The energy content varied still less: \pm 1.1 per cent and \pm 0.6 per cent for the two sets of samples.

Variable Part of the Ration.—The reference substance was composed of glucose as a standard for "nitrogen-free" food energy, and of casein as a standard food protein.

In extensive preliminary studies, the cows had apparently preferred rations containing glucose to those containing pure starch. Glucose has the further advantage of being a normal constituent of blood, whereas starch must be changed to become available to the body. The feeding of glucose instead of starch therefore meant a step forward in the simplification—in a biochemical sense—of the diet. This simplification, to be sure, may be considerably less effective in ruminants than it would be in one-stomach animals, because in cattle a considerable part of the glucose may not be absorbed into the blood-stream as such, but may be fermented by the microflora in the rumen.

Glucose was used in the form of "Cerelose." The carbon content analyzed during the first series of trials amounted to only 38.8 per cent instead of 40. When the discrepancy was discovered, it was too late to check up on the samples used for the carbon determination. Since the product seemed rather uniform and pure, there are grounds for suspecting an error in the carbon analysis by wet combustion. The error thus possibly created amounts to 1.4 per cent of the digested carbon in trial 16. The results were calculated on the basis of the theoretical carbon content of glucose.

Since casein represents one of the major parts of cow's milk, it can be assumed to have a high biological value, especially for lactation. Again the possible changes in the rumen may reduce the importance of this point.

The casein fed was the commercial product of 80 mesh. Table 3 gives its nitrogen, carbon, and energy content in comparison with the composition of pure casein according to Hammarsten (Richmond, 1920). The major impurity of the casein used in the trials was lactose.

In preliminary trials the cows ate the reference substance better when it was mixed with molasses beet pulp and was slightly moistened. Since poor appetite was the major difficulty, molasses beet pulp was included, and fed in about the same amount as casein. In comparing two production rations, one may count this molasses beet pulp together with the constant part of the ration. Table 3 gives the carbon, nitrogen, and energy content of the molasses beet pulp used.

The barley whose casein-glucose equivalent was measured in the trials was Western No. 1, weighing about 46 pounds to the bushel (59 kg per hectoliter). A sample contained (per 100 grams of dry matter) 5.8 grams of ash, 9.4 of protein, 1.5 of fat, 7.09 of crude fiber, and 76.3 of nitrogen-free extract.

TABLE 3

AVERAGE NITROGEN, CARBON, AND ENERGY CONTENTS OF FEEDS USED IN THE

MAIN RESPIRATION TRIALS

In 100 grams dry matter of	Trial series	Nitrogen	Carbon	Energy
		grams	grams	kcal*
Sudan hay	1	1.98	42.8	421
Sudan hay	2	2.24	43.2	428
Molasses beet pulp	1	1.16	42.7	403
Molasses beet pulp	2	1.23	43.8	413
Casein (commercial)	1	13.58	49.7	545
Casein (commercial)	2	14.84	52.8	571
Barley	1	1.65	43.2	435
Barley	2	1.94	44.3	441
Casein, pure (Hammarsten)		15 65	53 0	586
Glucose, pure C ₆ H ₁₂ O ₆			40.0	374

^{*} See footnote 11, p. 528.

To the ration was added bone meal amounting to 3 per cent and common salt amounting to 1 per cent of the dry reference substance.

Weigh-backs.—Whenever the cow left some food in the manger, these weigh-backs were collected, dried, and analyzed. In the last trial one cow left as much as 448 grams of dry matter per day. The approximate amounts of glucose, casein, and beet pulp in the weigh-back were estimated by means of chemical analysis.

Table 4 summarizes the rations ingested daily.

RESPIRATION TRIALS

Changes in the Respiration Apparatus.—The respiration apparatus used, the experimental procedure, and the method of calculating the results have been described in detail (Kleiber, 1935). Some improvements have been made. The reversing switch for moving the aspirator pipettes up and down has been replaced by an eccentric driving a rack and pinion; this made the operation quieter and more reliable. Instead of the water valves originally installed on top of the aspirator pipettes, the machine now operates with big ground-in steel stopcocks, which are greased with no. 3 cup grease mixed with graphite, and which work satisfactorily and smoothly.

To maintain a slight suction inside the chamber during the "dead period" of

TABLE 4
RATIONS CONSUMED DAILY, AS DRY MATTER

	RATIONS C	ONSUMED	DAILY,	AS DRY	MALIER			
	Level of food intake	Cow no.	Body weight, kg	Consta of rat	nt part	Variable part of ration		
Trial no.				Sudan hay, grams	Molasses beet pulp, grams	Casein, grams	Glucose, grams	Barley, grams
~ ~~	·	First se	ries of tria	ıls				
		494	467	5,270				
3	Maintenance	1003	493	5,370		• • •		
5	Maintenance	{ 494	488	4,455		***		
Ð	manitenance	1003	501	4,455				
7	Maintenance	∫ 1009	534	5,715				
•	3.300.200.100.100.100.100.100.100.100.100.1	1003	500	5,312				
9	Maintenance	∫ 1009	540	6,100				·
Ü		1003	517	5,680				
12	Lactation	∫ 1007	463	5,296	530	529	2,109	
		1021	474	4,883	530	529	2,109	
14	Lactation	∫ 1007	463	5,236	519	525	2,109	: : : : :
		1021	478	5,126	475			3,281
16	Lactation	∫ 1007	460	5,258	519	521 -	2,110	
		1021	473	5,258	475	318		3,118
18	Lactation	∫ 1007	470	5,219	386	241		2,393
		1021	473	5,219	386	402	1,646	
20	Lactation	∫ 1007	464	5,274	395	402	1,646	
		\ 1021	473	5,274	395	402	1,646	
22	Maintenance	∫ 1007	457	5,318				
		1021	466	5,318				
		Second s	eries of tr	ials		·	· · · · · · · · · · · · · · · · · · ·	
		∫ 732	409	4,257				
24	Maintenance	728	372	2,151				
		732	432	2,186				
25	Maintenance	728	371	3,243				
		∫ 732	434	2,186		144		
26	Maintenance	728	380	2,186		144	1,107	
		732	430	2,186		144		
27	Maintenance	728	382	2,186		144 144	1,107	
		732	378	3,580	400	376	3.232	
28	Lactation	728	347	3,580	400	90	3,232	4,183
-	T	732	373	3,580	400	90		4 100
29	Lactation	728	347	3,580	327	308	2,981	4,183
		L			1			

the aspirator, there was installed on top of each chamber a bellows made of an automobile inner tube slit open lengthwise and connected to the top of the chamber and to a circular sheet of aluminum. The center of this sheet is connected to a steel spring, which in turn is suspended from the ceiling of the room. The spring is made of steel wire, 2 mm thick, wound to a diameter of 2.5 cm; and a length of 1 m (in unstretched condition) is used for each bellows. A load of 1 kg stretches the spring 31 cm. When the aspirator sucks air out of the chamber, the metal plate of the bellows is drawn down to the roof of the chamber, since the inside of the bellows is connected to the inside of the chamber. When the aspirator reaches its dead period and there is no suction, the spring draws the metal sheet away from the chamber, thus drawing air from the chamber to the bellows. According to a test with a sensitive diaphragm manometer connected to a kymograph, this action of the bellows prevents a complete loss of suction in the chamber during the dead period of the aspirator.

The mercury in the sampling device becomes rapidly contaminated, probably with sulfide, when it pumps moist air that comes from the animal chamber. This trouble is almost completely avoided when the air is dried before coming in contact with the mercury. To eliminate weekly cleaning of the mercury in the sampling device, the air sample is dried before it reaches the mercury pump. Necessarily, however, this drying introduces considerable dead space. To avoid the resultant errors, the aspirator now pumps for every cycle (500 liters of ventilation air) a sample of 1 liter through a drier with phosphorus pentoxide and then back to the air line. From this sample of the first order, the mercury pump takes the secondary or actually used sample of 20 ml per cycle for the absorption battery and of 0.5 ml per cycle for the collector of the composite sample, which is later analyzed in a Haldane apparatus modified by Kleiber (1933) and mechanized by Winchester (1938).

Environmental Conditions.—The chambers were kept at about 20° C and at a relative humidity of 60 per cent.

RESULTS

DIGESTIBILITY OF FEED CONSTITUENTS

Digestibility of Nitrogen and Energy in Sudan Hay.—The digestibility as

used here is the quotient $\frac{\text{food minus feces}}{\text{food}} \times 100$. This is often called "apparent

digestibility" because part of the substance excreted in the feces is not undigested residues of the ingested food, but is material that has passed the metabolic process of the animals and is excreted into the intestinal tract.

In the first series, the digestibility was measured during the preliminary period of each experiment in order to check upon the results of the relatively short periods. On the average, the digestion coefficients in preliminary trials were equal to those of the main periods (table 5).

Statistically the slight decrease of the digestibility in the second series (61.7 + 1.0) compared with that in the first (62.5 + 0.9) is insignificant. The mean digestibility of uitrogen in Sudan hay for fourteen main trials on

maintenance is 61.9 ± 1.8 per cent. The digestibility of energy seems considerably higher for the Sudan hay used in these trials than the quotient

total digestible nutrients × 100, which for Sudan hay before bloom (Morrison,

1936) amounts to 55 per cent. In our trials, the digestion coefficient for energy in Sudan hay was 67.8 ± 0.3 and 66.2 ± 0.9 per cent, first and second series

TABLE 5

DIGESTIBILITY OF NITROGEN AND ENERGY IN SUDAN HAY

The state of the s	Number	Average d	igestibility
Experiments	determi- nations	Nitrogen	Energy
	10	per cent	per cent 67.7+0.5
First series, preliminary respiration trials	10	62.1±1.5	
First series, main respiration trials	10	62.5±0.9	67.8±0.3
Second series, main respiration trials	4	61.7±1.0	66.2±0.9
			1

TABLE 6
TOTAL AND PARTIAL DIGESTIBILITY OF ENERGY IN SUDAN HAY

		Total digestibility			Partial digestibility		
Cow no.	Trial no.	Food energy consumed per day	Energy digested per day	Digesti- bility	Change in food energy consumed per day	Change in energy digested per day	Partial digestibility
		kcal*	kcal	per cent	kcal	kcal	per cent
728	{ 24	9,290 13,982	6,288 9,006	67.7 64.4	4,692	2,718	57.9
732	{ 25	9,393 18,418	6,152 12,356	65.5 67.1	9,025	6,204	68.7

^{*} See footnote 11, p. 528.

respectively, or 67.2 ± 0.4 per cent for all fourteen main trials, when the results are statistically weighted for the duration of the trials.

In the second series, two trials were conducted to compare the utilization of Sudan hay fed at the half maintenance and at the maintenance level. Table 6 shows the digestion coefficients for energy that resulted from these trials; it serves especially to illustrate the meaning and the calculation of partial in contrast to total digestibility.

The term partial digestibility is chosen somewhat in analogy to the terminology used in basic sciences. The term "partial quantity," to be sure, is reserved in chemistry (Lewis and Randall, 1923, p. 34) for partial differential quotients, whereas quotients of partial differences are called apparent quantities. "Apparent" is, however, a rather poor adjective in this connection, because what "appears" is always a total, not a partial, effect (though partial effects may in special cases closely approach total effects). Aside from this criticism of "apparent," the term "apparent digestibility" is already used in

the literature. All the digestibility figures in the present paper are results for apparent digestibility (as distinguished from the "true" digestibility obtained by subtracting the metabolic nitrogen from the total nitrogen in the feces). The term apparent digestibility can therefore not be used to distin-

guish the quotient
$$\frac{\Delta \text{ digested}}{\Delta \text{ food}}$$
 from $\frac{\text{digested}}{\text{food}}$.

When at some future time the relation of digestibility to level of food intake

is known accurately enough, one may have to call $\frac{\Delta \ digested}{\Delta \ food}$ the "mean partial

digestibility" over a given range of food intake, in order to reserve "partial

digestibility" for the partial differential quotient $\frac{\delta \text{ digested}}{\delta \text{ food}}$.

Partial Digestibility of Nitrogen in Casein and Barley. The following example (based on trial 16, cow 1007) shows how the partial digestibility of protein in casein was calculated:

SAMPLE CALCULATION 1: PARTIAL DIGESTIBILITY OF NITROGEN IN CASEIN

1. Total nitrogen digested per day:

2. Partition of nitrogen digested per day:

3. Partial digestibility of nitrogen from casein:

In the first series, the partial digestibility of casein was 71 ± 1.5 per cent for cow 1007 (trials 12, 14, 16, and 20) and 72 ± 0 for cow 1021 (trials 12, 18, and 20). In the second series it was much more variable and lower—namely, 51 per cent for cow 732 in trial 28 and only 36 per cent for cow 728 in trial 29. During maintenance trials the partial digestibility of casein was highly variable—as might be expected, since the amounts fed (21.49 grams nitrogen in casein per day) were less than one half of those fed during the production trials. When casein and glucose were fed alone in the morning and Sudan hay alone in the evening, the partial digestibility of casein nitrogen was 56 per cent for cow 728 (trial 27) and as high as 71 per cent for cow 732 (trial 26).

The mean of the partial digestibility of nitrogen in case in the nine lactation trials of both series is 65.2 ± 4.3 per cent.

¹⁰ In the sample calculations, figures in **bold** faced type represent major results. Figure in ordinary type represent intermediate calculations.

The partial digestibility of nitrogen from barley was 67, 88, and 74 per cent for the first series. The low value of 67 resulted, in trial 14, with the low protein content of the ration having no casein supplement. During the second series of trials, the partial digestibility of nitrogen in casein was 67 and 64 per cent for cows 732 and 728 respectively. The mean of the partial digestibility of nitrogen in barley in five lactation trials is 72 ± 4.3 per cent. According to Watson and his co-workers (1939), the partial digestibility of nitrogen in barley, fed together with timothy hay, is 75.1 per cent. (Morrison, 1936, gives for eastern barley a protein digestibility of 79 per cent.) The difference between the nitrogen digestibility in casein and barley is not statistically significant.

According to Mendel and Fine (1912), meat did not increase the fecal nitrogen exerction of dogs over that resulting from a nitrogen-free diet, an observation confirmed by Mitchell and Carman (1924) on the rat with egg and pork added to a nitrogen-free diet. That in the cows casein should be no more digestible than barley protein may therefore seem surprising.

One may assume that the nitrogen from casein is actually fully digestible, and regard the low partial nitrogen digestibility of the rations with the reference substance as the result of a depression of the apparent nitrogen digestion by glucose. That carbohydrates depress the digestibility of protein was indeed suggested as early as 1870 by Stohmann (Mayer, 1925), who gave a formula to calculate the apparent digestibility of protein as a function of the carbohydrate content of the ration. G. Kühn (Kellner, 1919, p. 53) observed that starch decreased the digestibility of hay protein from 57 to 49 per cent when 1.67 kg of starch was added, and to 42 per cent when 2.87 kg of starch was added, to the same amount of hay. Armsby (1922, p. 618) concludes from Pfeiffer's and Kellner's data that "the digestion of each 100 grams of dry matter, whether protein or nitrogen-free material, results in the excretion in the feces of approximately 0.4 gram of nitrogen in the form of excretory products," which products he regards (p. 109) as composed of "unresorbed digestive juices and their decomposition products, intestinal mucus, worn-out epithelial cells and cell fragments, leucocytes and excretions of the intestinal mucosa." An increase of this intestinal excretion. according to Armsby, causes the depression of the apparent nitrogen digestibility by carbohydrate feeding.

The nitrogen in these excretory products was later designated as metabolic nitrogen. Mitchell (1926, p. 34), on the basis of experiments of Morgan, Berger, and Westhauser on sheep, assumes that the excretion of metabolic nitrogen, per 100 grams of dry matter consumed, amounts to 0.5 gram when the rations contain considerable amounts of crude fiber. "It seems," writes Mitchell, "that the differences in apparent digestibility observed may be explained entirely on the demonstrated property of carbonaceous food of causing an excretion of metabolic nitrogen in the feces in proportion to its dry matter content."

Hamilton (1942, p. 107) noted, in trials with sheep, that glucose, added in amounts of 20 to 30 per cent to a ration of timothy hay, yellow corn, and cottonseed meal, reduced the digestibility of nitrogen from 62 to 54 per cent. This depression of nitrogen digestibility can be accounted for, according to

Hamilton, by an estimated increase in the metabolic nitrogen in the feces of the sheep on the sugar ration.

The concept of metabolic nitrogen, however, does not seem to be sufficiently clear yet as a basis for a satisfactory explanation. On page 21 Mitchell (1926) states that "a certain fraction of the fecal nitrogen, the so-called 'metabolic nitrogen' of the feces, is contained in substances originating in the animal body." On page 27 he includes in the metabolic nitrogen of the feces also the "waste nitrogen from digestion," in addition to the waste nitrogen from the body. On page 29 he writes: "It appears, therefore, that the so-called 'metabolic' nitrogen of the feces consists predominantly of nitrogenous waste products of digestion rather than metabolism."

Why a soluble nitrogen-free substance increases nitrogenous waste products of digestion, is a problem that needs further clarification.

Aside from nitrogen digestibility, the addition of glucose distinctly depressed also the digestibility of crude fiber in steer calves (Mitchell, Hamilton, and Haines, 1940) and in sheep (Hamilton, 1942). An explanation of the depression of nitrogen digestibility in ruminants may be attempted on the basis of these observations. If less fiber is digested with the added sugar, presumably more food protein remains protected by fiber against the action of digestive enzymes and absorption through the intestinal tract. In this case, however, the depression would involve the "true" digestibility; unabsorbed food nitrogen, rather than metabolic nitrogen, should be increased in the feces. One does not have to assume with Hamilton that rumen bacteria (like people) "prefer" sugar to fiber; sugar may, to the contrary, have an adverse effect on the rumen flora, or at least on that part of this flora which attacks fiber. Any reduction of bacterial activity could account for decreased fiber digestion. A suitable antiseptic in the rumen would probably lead to a decrease in fiber digestion. Following his reasoning in the Journal of Nutrition (based solely on digestion data), Hamilton would then have to conclude that bacteria which, in the absence of the bactericide, attack cellulose will prefer the bactericide to the fiber.

Woodman and Evans (1938) have demonstrated that rumen bacteria in vitro ferment glucose readily. Possibly the split products of glucose resulting from this fermentation inhibit or at least depress the fermentation of fiber. An increase in fecal nitrogen may of course also result from an increase in bacterial growth possibly favored by sugar and leading to an increase in bacterial protein in the feces.

The depression of the apparent digestibility of protein by carbohydrates is not yet satisfactorily explained. Our trials do not lead to the solution of this problem either. We therefore simply report the results as partial effects, reserving possible explanations to later investigations.

The partial digestibility of the nitrogen in the casein, namely 72 per cent for the first series of the trials, indicates that the fecal nitrogen was increased 0.28 gram for each gram of nitrogen in casein. The reference substance contained 25 grams casein or 3.4 grams nitrogen per 125 grams of dry reference substance or per 100 grams glucose. The partial fecal nitrogen resulting from

the reference substance thus amounts to $3.4 \times \frac{100}{125} \times 0.28 = 0.78$ gram nitrogen

per 100 grams of dry reference substance or to $3.4 \times 0.28 = 0.95$ gram nitrogen per 100 grams of glucose. This is about twice the effect observed in the earlier trials reported above. For the second series, with only two results and much

more variability, the depression is still greater.

Partial Digestibility of Energy in Glucose and Barley.—Sample calculation 2 (based on trial 16, cow 1007) shows how the partial digestibility of energy from glucose was calculated. The mean for this partial digestibility, calculated from seven main production trials, is 83 ± 1 per cent. This partial digestibility was higher in the second series—89 and 88 per cent, respectively, for cows 732 and 728. Still somewhat higher, but not significantly different from the last results, are those from maintenance trials in the second series, with a mean for four trials of 93.5 ± 3.4 per cent.

Sample Calculation 2: Partial Digestibility of Energy in Glucose

1. Total energy digested per day:	•
Energy in feed per day	35,150 kcal ¹¹
Energy in feces per day	10,110 kcal
Digested energy per day (35,150 — 10,110)	
O. Destition of an area of a second and a second as a	

2. Partition of energy digested per day:

Energy digested from molasses beet pulp per day: Energy consumed per day in 519 grams dry molasses beet pulp is 2,095 kcal; digestibility (according to Morrison's 1936 tables) is 85 per cent; therefore energy digested from molasses beet pulp is $0.85 \times 2,095...1,781$ kcal

3: Partial digestibility of energy in glucose:

Partial digestibility of energy in glucose: $\frac{6,571}{7,885} \times 100.$ 83 per cent

These results do not imply that 10 to 20 per cent of the glucose was excreted in the feces. Rather, they mean that the addition of glucose slightly lowered the digestibility of food energy. This depression is superimposed upon that of protein digestibility, whose effect on energy digestion is already accounted for in the calculation of the partial digestibility of glucose (sample calculation 2), since only the partially digested fraction of the casein energy figures in that calculation. The partial digestibility of the energy in the reference substance as a whole (casein and glucose) amounts to 80.0 ± 0.7 per cent. The addition of glucose distinctly depressed the digestibility of crude fiber in steer calves (Mitchell, Hamilton, and Haines, 1940) and in sheep (Hamilton, 1942).

The partial digestibility of energy in barley, calculated like that for glucose (sample calculation 2), was 74 per cent for the trial with the low protein con-

[&]quot;In this paper, kilocalorie will be abbreviated kcal. A kilocalorie is 1,000 calories. The term "large calorie" is obsolescent, and the term "kilogramcalorie" is a misnomer.

tent and 80 per cent for the diet that was balanced by casein (cow 1021). Since it was 74 per cent for the balanced diet with cow 1007, the difference with cow 1021 may be taken to result from variability rather than balance of the diet. The partial digestibility of barley energy in the second series was 78 and 77 per cent respectively. The four trials with balanced rations led to a mean (weighted with respect to duration of the trials) of 77.3 ± 1.1 per cent partial digestibility of the energy in barley. From Morrison's table (1936), by way of comparison, one may calculate for eastern barley an energy digesti-

bility
$$\left(\frac{\text{total digestible nutrients}}{\text{organic matter plus } 1.25 \times \text{fat}}\right)^{12}$$
 of 87 per cent.

METHANE

Methane Production and Intake of Sudan Hay.—In the first series of experiments the methane production was rather uniform. The mean for five main trials (ten results) on maintenance with Sudan hay alone was 37.0 ± 0.7 liters methane per kilogram dry Sudan hay eaten. Since 1 liter methane weighs 0.715 gram and since 100 grams of dry Sudan hay contains 75.4 grams of carboyhydrates (p. 520), the methane production in the first series of the trials was 3.5 ± 0.07 grams methane per 100 grams of carbohydrates in Sudan hay.

Using our analysis of the Sudan hay and measurement of digested nitrogen, and assuming 40 per cent digestibility for fat (Morrison, 1936), one can estimate that the digestibility of the carbohydrates in Sudan hay was 79 per

cent. The methane production, consequently, amounted to
$$\frac{3.5}{0.79} = 4.4$$
 grams

methane per 100 grams of digestible carbohydrates in Sudan hay. This confirms Kellner's results on steers, as summarized to 4.29 grams of methane per 100 grams of digested carbohydrates (Kellner, 1919, p. 96), and those of Armsby and Fries (1915, p. 450) with 4.8 grams of methane per 100 grams of digested carbohydrates as a mean of 43 experiments with roughages, and 4.7 grams of methane as a mean of 22 experiments with rations mixed of roughages and concentrates. Later calculations at Pennsylvania State College (linear interpolations by the method of least squares) produced essentially the same result, adding a complicating rather than clarifying term that implies methane production without food intake (Kriss, 1930; Bratzler and Forbes, 1940).

The methane production during the second series of trials was more variable than during the first. On full-maintenance rations of Sudan hay cow 732 produced 35 and cow 728 produced 40 liters of methane per kilogram of dry

hay. The latter's partial methane production
$$\left(\frac{\text{change in methane}}{\text{change in hay}}\right)$$
 between

half maintenance and maintenance was 37 liters per kilogram of dry Sudan hay. Cow 732, however, produced 51 liters of methane per kilogram of dry hay on half-maintenance ration; and her partial methane production between half maintenance and maintenance on Sudan hay was as low as 18 liters of methane

¹² Fat-free organic matter plus 2.25 × fat, or organic matter plus 1.25 × fat.

per kilogram of dry Sudan hay. Aside from that one trial on half maintenance with cow 732, our results in the second series of trials confirm those in the first.

Partial Methane Production Resulting from Reference Substance.—Sample calculation 3 (based on trial 16, cow 1007) illustrates the calculation of partial methane production resulting from glucose, on the assumption that casein does not contribute to methane formation. This assumption seems reasonable, since Kellner and his co-workers stated as early as 1896 (p. 287) that they could not detect any influence of protein on the methane formation of steers.

SAMPLE CALCULATION 3: PARTIAL METHANE PRODUCTION FROM GLUCOSE

1. Total methane production per day:

2. Partition of methane production:

3. Partial methane production per 100 grams glucose:

Consumed per day: 2.110 kg glucose (dry matter). Methane production per 100 grams

The mean partial methane production from glucose, calculated from seven main trials with two cows during the first series of the experiments, amounted to 1.8 ± 0.34 grams of methane per 100 grams of dry glucose. This is only half the methane production resulting from the carbohydrates in hay $(3.5\pm0.07$ grams of methane), as mentioned above; and the difference is statistically highly significant. In the second series, however, the partial methane production from glucose is higher—namely, 2.4 grams and 2.6 grams of methane per 100 grams of glucose for cows 732 and 728 respectively. In experiments with steers, Mitchell, Hamilton, and Haines (1940, p. 857) did not observe any increase of methane production resulting from the addition of glucose to a basic diet.

With the maintenance rations of the second series, the methane production per 100 grams of glucose was higher than with the production rations. It ranged from 2.9 to as high as 7.4 grams of methane per 100 grams of glucose. The greater variability, the smaller number of trials, and the lower food intake tend to make the last-mentioned results less reliable than those of the first series. A distinct increase of the methane fermentation (per 100 grams of carbohydrate) with a decrease of the rations (alfalfa hay and starch) has been reported by Armsby and Fries (1918, p. 274).

Partial Methane Production Resulting from Barley.—The partial methane production from barley in the first series of trials was 3.6 and 3.2 grams of methane per 100 grams of carbohydrates in barley for cow 1021 and cow 1007 respectively. The corresponding figures for the two cows in the last series are

2.1 and 4.1 grams of methane, so that the mean for the four main trials is 3.2 ± 0.4 grams of methane per 100 grams of carbohydrate in barley, a figure not significantly different from that for carbohydrates in Sudan hay $(3.5\pm0.07~{\rm grams})$.

METABOLIZABLE ENERGY

Definition.—Metabolizable energy is that part of the total food energy which is available for any metabolic process in the animal—namely, katabolism when chemical energy is transferred to work or heat; and anabolism, whereby the chemical energy of food is transferred to the chemical energy of animal products—for example body fat, body protein, milk, eggs, or wool. The metabolizable energy is the difference between the heat of combustion in the food and that in all excreta (feces, urine, and methane).

The heat produced by katabolism in the animal body from the chemical energy of the food is regarded here as part of the metabolizable energy of the food, whether it is evolved by the metabolism of microörganisms in symbiosis or parasitosis with the animal or by the metabolism of the animal's own cells. The heat increment likewise includes increase in heat production by increased metabolism of body cells or by increased metabolism of microörganisms in the animal body. From the point of view of energy exchange we are used to regarding the animal, together with its symbionts and parasites, as a unit. Excluding the heat produced by the microörganisms in the animal body from the metabolizable energy would mean, at the present state of research, defining metabolizable energy as a hypothetical rather than a measured quantity.

Metabolizable Energy in Sudan Hay.—The metabolizable energy of Sudan hay in ten main trials on the maintenance level with dry cows averaged to 222.5 \pm 1.4 kcal per 100 grams of dry matter. The variability is rather low, the coefficient of variation being \pm 1.9 per cent. Even slightly lower—namely, \pm 1.7 per cent—is the coefficient of variation for the metabolizability of the digested energy in Sudan hay (metabolized energy/digested energy), which, as a mean of ten results in the main trails on maintenance, is 77.8 \pm 0.4 per cent. If the preliminary trials are included, then the mean amounts to 77.4 \pm 0.4 per cent. In two maintenance trials of the second series of the experiments 100 grams of dry Sudan hay contained 228 and 215 kcal of metabolizable energy. The details are found in table A of the appendix.

Partial Metabolizable Energy in Reference Substance.—Sample calculation 4 (based on trial 16, cow 1007) illustrates the calculation of the partial metabolizable energy in the reference substance. The metabolizable energy in molasses beet pulp has been estimated from the data in Morrison's tables (1936) and from the mean of 1,616 kcal of metabolizable energy per pound of total digestible nutrients given by M. Kriss (1931). Such an estimate was justified because molasses beet pulp furnished only 7 per cent of the metabolizable energy in the rations.

Sample Calculation 4: Partial Metabolizable Energy in Reference Substance and in Glucose

- 2. Partition of metabolizable energy per day:

Metabolizable energy from Sudan hay: The Sudan hay used contained 2.22 kcal of

casein and 2,110 grams glucose.

3. Partial metabolizability of energy in reference substance:

bility of energy in glucose amounted to $\frac{5,810}{7,885} \times 100...$ 74 per cent

As a mean of seven trials with two cows in the first series of experiments, the partial metabolizable energy in the reference substance was 301 ± 5 keal per 100 grams dry reference substance, or 74 ± 1.3 per cent of the total energy in the reference substance.

The utilization of nitrogen from casein was very high, as will be shown later. (See p. 540.) If we assume that the digested energy from casein was 100 per cent metabolizable (no increase in urinary nitrogen resulting from feeding casein), then we calculate the partial metabolizable energy in glucose alone as shown in sample calculation 4. According to this calculation for the seven main production trials, the mean partial metabolizable energy in glucose was 278 ± 6.3 kcal per 100 grams of glucose, or 75 ± 1.8 per cent of the total energy in the glucose. In the second trial the metabolizability of glucose seemed somewhat higher—namely, 298 and 290 kcal of metabolizable energy per 100 grams glucose, or 80 and 78 per cent of the energy in glucose for cows 732 and 728 respectively.

The duration-weighted means for both series (20 weeks of trials) of partial metabolizable energy in glucose are 283 ± 5 kcal metabolizable energy per 100 grams glucose, or 76 ± 1.5 per cent of the total energy in glucose.

The metabolizability of glucose for maintenance, according to a similar calculation, was rather variable: as a mean of four trials it amounted to 292 ± 24 kcal per 100 grams of dry glucose, or 78 ± 6 per cent of the total energy in glucose.

Mitchell, Hamilton, and Haines (1940) report for steer calves a metabolizable energy in glucose averaging to 65 per cent of the heat of combustion of glucose. They attribute (p. 861) this low metabolizability to the partial de-

struction of glucose by the paunch flora and to the depressing effect of glucose on the digestibility of polysaccharides.

The partial destruction of glucose in the rumen can, however, hardly be responsible for a low metabolizability, since Mitchell, Hamilton, and Haines found no increase in methane production resulting from the addition of glucose to the ration; the only possible way in which this partial destruction could otherwise account for a lowering of the metabolizability—as this term is defined here and as it is calculated by Mitchell himself—would be the formation of split products excreted in the urine.

Partial Metabolizable Energy in Barley.—The partial metabolizable energy in barley can be calculated from three main production trials of 2 weeks' duration and two trials of 3 weeks' duration by procedure analogous to sample calculation 5. The duration-weighted means resulting from such calculations are 293 ± 9.7 kcal metabolizable energy per 100 grams of dry barley and a metabolizability of 67 ± 2 per cent of the total energy in barley.

MAINTENANCE REQUIREMENT AND SUDAN HAY AS FOOD FOR MAINTENANCE

Definition.—Maintenance means a condition in which an animal is in nutritive equilibrium, which is attained when it neither gains nor loses. Gain or loss may concern any nutritive essential. Thus we may speak of energy maintenance when the animal neither gains nor loses energy; or protein maintenance when no gain nor loss of nitrogen from the body occurs. In a similar way we can speak of calcium maintenance, phosphorus maintenance, even water maintenance. Maintenance is a dynamic equilibrium. An animal on protein maintenance, for example, may actually lose nitrogen that was part of its tissues; but it replaces this loss by the same amount of nitrogen retained from its food. Maintenance is thus defined as the condition under which the intake of a certain nutritive essential is equal to the sum of all losses of that essential. A fasting animal obviously cannot be on maintenance, according to our definition, which thus differs from the concept of Forbes and his coworkers (1941, p. 19), who apparently use the word maintenance in a wider sense, such as keeping alive, whereas we restrict it to a definite quantitative term.

Maintenance requirement is the amount of a certain food that provides maintenance. Even for a given food, this requirement will of course usually differ for different nutritive essentials. Though a certain amount of Sudan hay may suffice to keep an animal in calcium equilibrium, this amount may be inadequate to keep it in phosphorus equilibrium; more hay was necessary in our trials to provide energy equilibrium than to provide protein equilibrium. (See p. 570.) Appendix table B summarizes results on feed utilization.

The maintenance requirement changes with the animal's size; and the function between size and maintenance requirement may be different for different essentials. Energy requirement, for example, is closely proportional to the 34 power of body weight (Kleiber, 1933), whereas the vitamin-A requirement is proportional to the body weight itself (Guilbert and Hart, 1935).

In this study the requirements of protein and energy only were considered. Fortunately, these two requirements are related to the same function of looky

size, as was discovered by Terroine and his co-workers (1927) and called the Terroine, Sorg-Matter law. This law was confirmed by D. B. Smuts (1935, p. 427), who summarized his own results, with those of Terroine et al., in the statement that 2 mg of nitrogen is required for each "calorie" (in our terminology, kilocalorie, kcal) of basal heat.12

Since basal-heat production is most nearly proportional to the 3/4 power of body weight (Kleiber, 1932),14 the rate of food intake, as well as the rate of gain and loss of protein or energy in body substance, was calculated per unit of the 3/4 power of body weight. Thus was eliminated as much as possible the effect of differences in body size of the cows. This procedure has been justified against criticism in an earlier publication (Kleiber, 1938). Table C of the appendix gives the metabolic size for cows of 600 to 1,400 pounds' body weight.

Requirement of Sudan Hay for Protein Maintenance .- In all eight of the main maintenance trials with Sudan hav carried out in the first series of experiments, the cows showed slight gains of nitrogen. There was no correlation between these gains and the nitrogen intake. To get information on the utilization of nitrogen from Sudan hay, we calculated the basic nitrogen requirement according to Smuts' rule mentioned above, using as daily rate of basal metabolism the result of our fasting trials—namely, 86 kcal per kg 4.15 (See p. 535.) This leads to a daily maintenance requirement of 0.172 grams net nitrogen per kg%.

Sample calculation 5 illustrates the calculation of protein utilization on this basis. For the eight trials of the first series, the mean net-nitrogen ontent in Sudan hay was 0.49 ± 0.03 gram of net nitrogen per 100 grams of dry matter, which is 25 ± 1.4 per cent of the total nitrogen and 40 ± 2.4 per cent of the digested nitrogen in Sudan hay.

SAMPLE CALCULATION 5: NET NITROGEN IN SUDAN HAY FOR MAINTENANCE; AND REQUIRE-MENT FOR PROTEIN MAINTENANCE

1. Total net nitrogen per day:

With a basic nitrogen requirement of 0.172 gram net nitrogen per kg% (as explained above), cow 1007, trial 22, with a metabolic body size of 99 kg 34 , required daily 99 \times 0.172 or 17 grams net nitrogen. In her gained body substance she stored daily 13 grams nitrogen. Basic requirement added to gain equals total net nitrogen $17 + 13 \dots 30$ grams

2. Net nitrogen content of Sudan hay:

The 30 grams net nitrogen was supplied by the daily ration of 5,318 grams dry matter in Sudan hay containing 104 grams total nitrogen, of which 64 grams were digested. The Sudan hay therefore supplied net nitrogen as follows:

13 The creatinine fraction of the endogenous urinary nitrogen, however, seems to be in

proportion to body weight itself (Brody, Procter, and Ashworth, 1934).

14 Brody and Procter (1932, p. 99) in a later publication of the same year concluded that "the basal metabolism of mature mammals and of the domestic fowl increases directly with the 0.734 power of body weight." We do not believe that the second, let alone the third. decimal of the exponent has at present real significance; the difference between Brody's 0.734 and of Kleiber's more rounded 34 power appears to be unimportant.

 15 In this publication W^{3_4} means the body weight in kilograms raised to the % power, or the metabolic body size of the animal; kg% stands for the unit of this metabolic body size. If a metabolic rate (in terms of nitrogen or energy) is given per kg 4, one can estimate the rate of an animal of any size as: (rate per kg%) times metabolic size. The metabolic size of cows weighing 600 to 1,400 pounds is given in table C of the appendix.

¹⁶ See note p. 518.

3. Requirement for daily protein maintenance per unit of metabolic body size:

Dry matter in Sudan hay:
$$\frac{0.172}{0.56} \times 100 = 31$$
 grams per kg%
Total nitrogen in Sudan hay: $\frac{0.172}{29} \times 100 = 0.59$ gram per kg%
Digested nitrogen in Sudan hay: $\frac{0.172}{47} \times 100 = 0.37$ gram per kg%

The nitrogen losses in urine in one of four trials of the second series were far out of line with all the other corresponding data. If this result is omitted, a mean of 0.59 ± 0.03 gram net nitrogen per 100 grams of dry Sudan hay is found for the second series. This is 25 ± 1.5 per cent of the total nitrogen or 42 ± 2.6 per cent of the digestible nitrogen. Although the hay had a higher nitrogen content in the second series than in the first (see table 3), the utilization of the nitrogen for maintenance was the same in both series.

Sample calculation 5 shows how to calculate the requirement of Sudan hay for protein maintenance on the basis of Smuts' rule. This calculation, applied to the results of eight chief maintenance trials of the first series, results in a mean daily requirement of 36 ± 2 grams of dry Sudan hay, 0.71 ± 0.04 gram of total nitrogen or 0.44 ± 0.03 gram of digestible nitrogen per kg³4. The corresponding figures for the second series are 29 ± 2 grams of dry hay, 0.68 ± 0.04 gram of total nitrogen, or 0.41 ± 0.02 gram of digested nitrogen per kg³4.

Requirement of Sudan Hay for Energy Maintenance.—The daily fasting katabolism on the fourth and fifth day of fast was 89 and 83 keal per kg³⁴ for two of the cows respectively. These results are calculated on the basis of 4.7 keal per liter of O₂ consumed. It is considerably above the interspecific mean of 72 keal per kg³⁴ (Kleiber, 1932); or the mean of 71.75 keal per kg³⁴ reported by Mitchell, Hamilton, and Haines (1940) as the fasting heat production of four steers ranging in weight from 400 to 500 pounds; or the mean for fasting dry cows of 73 keal per kg³⁴ calculated from results given by Forbes and his co-workers (1927). The difference, which may be regarded as a stimulating effect of pregnancy on metabolic rate, is in line with our unpublished observations on rats. The energy loss (heat production of animal and chemical energy in urine) of the cows amounts to 93 and 90 keal per kg³⁴ of the two cows.

The basic requirement for net energy so far cannot be explained in terms of well-understood energy needs. The work of the heart and the respiratory system and even the osmotic work of internal organs do not seem to sum up to the actual energy expenses of a fasting and resting animal. Even if they did, it would be rather unsatisfactory to "explain" the oxidation of body tissue as necessary for the transport of oxygen, which in turn is necessary for the oxidation of body tissues. Apparently the only satisfactory teleological explanation of the basal metabolism is a comparison to the idling of a car, without starter, in order to have it ready for immediate action—that is, the mainte-

nance of a state of physiological alertness. At present, however, it is hard to explain rationally why, above the critical temperature, this idling should require a definite amount of energy, instead of some multiple or fraction of that amount. We can therefore reasonably talk of the heat requirement of a thermostat or of an animal below its critical temperature for maintaining a constant body temperature, but we have not yet a corresponding understanding for a requirement of energy other than heat, because we lack a criterion analogous to temperature. Before such a requirement is known, one can hardly speak of utilization of body substance by a starving animal (above the critical temperature), because utilization implies a purpose.

This consideration makes it somewhat difficult for us to understand the theoretical significance of Forbes and his co-workers' (1941) derivation of a "minimum base value of animal heat production." This term is defined as the difference between the basal metabolism and the heat increment, which latter would be observed if the body substance, katabolized by the fasting animal, were fed to the animal at or above the maintenance level. The derivation is based on the assumption that the animal has a certain minimum requirement of energy other than heat (but apparently kinetic) and that in order to meet this requirement a starving animal "utilizes" its own body substance, "wasting" some of the chemical energy of this substance as heat increment.

Even before an energy requirement (aside from heat) of a starving animal can be defined clearly, and before the heat increment of food is more completely understood, the "minimum base value of heat production" might be helpful as an empirical constant, useful for the presentation of animal heat production as a linear function of nutritive level. The minimum base value calculated by Forbes and his co-workers does not, however, seem to fit some of their own results such as those with different levels of intake of alfalfa hay and corn meal (1941, table 5, p. 22). Mitchell, Hamilton, and Haines (1940) question the validity of the experimental bases (as such) of Forbes's theory of the minimum base value.

Considering these difficulties, we decided that our calculations concerning the utilization of Sudan hay for energy maintenance should be based on the daily rate of fasting energy loss obtained by direct measurements. This is the simplest and clearest procedure, anyway. Sudan hay for energy maintenance is used, from the farmer's point of view, to prevent the losses of energy from body substance that would occur without food. These losses (heat loss of animal + chemical energy in urine) in our trials average 92 kcal per day per kg. Net energy in the maintenance food, or the saving of energy in body substance which would be lost without food, is thus 92 kcal per kg 4. None of our rations was exactly a maintenance ration. In all trials reported here, gain or loss of body substance and, consequently, of energy occurred. The total net energy of the ration is then calculated by adding the energy in the gained body substance to, or subtracting that in the lost body substance from, the net energy for maintenance, which was calculated as 92 times the metabolic size of the cow in kg. Sample calculation 6 demonstrates this calculation for cow 1007, trial 22, body weight 457 kg, metabolic body size 99 kg. There is a question whether the energy of gain is really directly comparable with the energy of loss-whether, in other words, partial efficiency of energy utilization above and below the maintenance level is equal. This question becomes, however, less important as the losses and gains of energy become smaller. It is not important when one succeeds in feeding close to the maintenance requirement.

SAMPLE CALCULATION 6: NET ENERGY IN SUDAN HAY FOR MAINTENANCE, AND REQUIREMENT FOR ENERGY MAINTENANCE

1. Total net energy per day:

During fasting, the cow (1007) lost daily as heat plus chemical energy in urine 92 kcal per kg. Her basic energy requirement in this trial (22) therefore amounted to 92 × 99, or 9,108 kcal. In her gained body substance she stored daily 870 kcal of chemical energy. Basic energy requirement added to gain in energy equals total net energy 9,108 plus 870

2. Net energy content of Sudan hay:

The 9,978 kcal net energy was supplied by the daily ration of 5,318 grams dry matter in Sudan hay containing 22,470 kcal total energy, of which 15,210 kcal was digested. The Sudan hay therefore supplied net energy as follows:

3. Requirement for daily energy maintenance per unit of metabolic body size:

Dry Sudan hay:
$$\frac{92}{188} \times 100 =$$
 49 grams per kg%

Total food energy in Sudan hay: $\frac{92}{44} \times 100 =$ 210 kcal per kg%

Digested energy in Sudan hay: $\frac{92}{66} \times 100 =$ 140 kcal per kg%

As an average of eight results in main trials of the first series, the net energy in Sudan hay for maintenance amounts to 176 ± 5 kcal per 100 grams of dry matter, 62 ± 2 per cent of the digested and 80 ± 2 per cent of the metabolizable energy in Sudan hay. The corresponding figures for the two trials of the second series are 135 and 151 kcal per 100 grams of dry matter, 47 and 54 per cent of the digestible and 59 and 70 per cent of the metabolizable energy.

When the net energy for maintenance in hay is known, then one can of course also estimate how much hay must be fed to produce neither gain nor loss of energy. Sample calculation 6 shows this procedure.

From the hay of the first series of trials the cows needed for energy maintenance (on the average of eight trials) 52 ± 2 grams of dry matter, 150 ± 4 keal of digested energy, or 116 ± 3 keal of metabolizable energy per day per kg*. The corresponding figures for the two maintenance trials of the second series are 68 and 61 grams of dry matter, 196 and 171 keal of digested energy, and 156 and 132 keal of metabolizable energy. Either the hay of the first series was definitely more nutritious for maintenance than that of the second series, or else the cows of the first series utilized the energy in the hay more efficiently than did the cows in the second series.

The mean requirement of Sudan hay for protein maintenance was 36 ± 2 grams per kg³⁴ per day in the first series; 29 ± 2 grams of dry matter per

kg" in the second. Thus the cows needed considerably more Sudan hay for energy maintenance than for protein maintenance.

The hay dry matter of the first series was more effective for energy maintenance, but less effective for protein maintenance, than the hay dry matter of the second series. Table 7 summarizes these results.

Sudan hay is more than rich enough in protein as a maintenance food.

Möllgaard's factor K, which is fattening energy in protein total fattening energy, of Sudan hay17

was 0.11, thus slightly higher than the 0.10 recommended by Möllgaard for maintenance. Judging from the present trials, a Möllgaard ratio of 0.08 in the maintenance ration would have provided sufficient protein.

TABLE 7 DRY MATTER OF SUDAN HAY NEEDED FOR MAINTENANCE PER DAY PER KG 34

Trial series	Protein main- tenance, grams per day per kg. ^{3/4}	Energy main- tenance, grams per day per kg. ^{3/4}
First (5 trials)	36±2 29±2	52±2 64±4

Partial Utilization of Reference Substance for Maintenance.—Four trials enable us to calculate the partial net nitrogen in casein for maintenance. One cow (732) seems to have been considerably and consistently more efficient in utilizing casein than the other cows. The means for four results are as follows: 100 grams of dry casein, added to a half-maintenance ration of Sudan hay, saved 16.7 ± 2.3 grams of nitrogen of the animal body (net nitrogen). The partial net nitrogen in casein under those circumstances amounted to 112 ± 15 per cent of its total nitrogen, or to 196 ± 16 per cent of its digestible nitrogen.

As mentioned before (p. 525), the partial digestibility of casein was rather low, especially in the maintenance trials. The added casein, however, increased as a supplement the biological value of the hay proteins correspondingly, so that 1 gram of casein added to the hay ration actually saved (on the average) 1 gram of body nitrogen from decomposition.

The same four trials just mentioned can be used to calculate the utilization of the energy in the reference substance (casein and glucose) for maintenance. The results vary a great deal. There is no significant difference in the energy utilization between the two trials during which easein and glucose only were fed in the morning and hay only in the evening, and the two other trials during which hay and reference substance were fed together twice a day. The reference substance (containing 11.5 per cent casein and 88.5 per cent glucose) had a partial net-energy effect for maintenance (in addition to Sudan hay) of 120 kcal net energy per 100 grams when fed together with the hay; and of 126 kcal net energy when fed alone in the morning, the hay being fed alone in the evening.

¹⁷ This hay contained 12.5 per cent protein, with a digestibility of 67 per cent.

The mean of all four results is 123 ± 35 kcal net energy per 100 grams of dry reference substance; or a partial efficiency for maintenance of 31 ± 10 per cent of the total, 36 ± 12 of the digestible, and 40 ± 9 per cent of the metabolizable energy in the reference substance.

An attempt to calculate the partial net energy in glucose alone, by assuming that the energy utilization in casein is equal to the nitrogen utilization in casein, led to a very low utilization of glucose—in one case even to a negative net-energy value of glucose for maintenance. The type of feeding—hay and reference substance mixed, or hay only in the evening and reference substance alone in the morning—did not influence the partial utilization of glucose, which on the average was 65 ± 39 kcal energy per 100 grams of glucose, corresponding to 18 ± 10 kcal of net energy per 100 kcal of total energy in glucose.

PROTEIN AND ENERGY UTILIZATION FOR LACTATION

Definition.—By protein utilization or energy utilization we mean in this publication the transfer of food protein or food energy to protein or energy in the animal's body or in the animal's product—that is, into a form which is useful from the farmer's point of view. We thus use the concept "utilization" in a more restricted sense than Forbes and his collaborators (1941): they mention the energy expense of utilization of katabolized body nutrients and the net energy of internal work, by which they probably mean the work of the heart and other organs necessary for providing oxygen for the katabolism of body nutrients. Since the farmer as a rule cannot use the heat produced by his animals, the burning up of body nutrients for internal work is, in animal production, not a utilization but a waste. The usefulness of the maintenance food is then measured by the amount of this waste that it prevents. This is the basis for Armsby's definition of the net energy for maintenance (1922, p. 271). The net energy of a ration for maintenance is the amount of energy of body substance that the ration saves from being transferred to heat.

The entire ration of a producing animal is useful in two ways: first, it prevents loss of body substance; second, it provides material and energy for the animal's product, such as milk, and for its gain in body substance. The net energy of such a ration can thus be formulated as follows: net energy of ration equals net energy for maintenance plus energy in product (milk, etc.) plus energy in gained body substance. This formulation is correct also when the animal loses body substance, because such a loss enters into the calculation as a negative gain.

In analogy to net energy, we may speak of net nitrogen as a criterion for the utilization of protein and may formulate: net nitrogen of ration equals net nitrogen for maintenance plus nitrogen in product plus nitrogen gain in body.

Partial Utilization of Nitrogen from Casein.—The net nitrogen for maintenance of the cows can be calculated on the basis of Smuts' rule mentioned before (p. 534), as follows:

N(net) for maintenance = 0.172 W^* , where W is the body weight of the cows in kilograms and N(net) is the number of grams of net nitrogen required per day. The calculation is based on the mean basal metabolism of 86 keal per kg* on the fourth and fifth day of fast.

Addition of the nitrogen in milk per day and the nitrogen gained daily as body substance to the net nitrogen for maintenance leads to the total net nitrogen per day of our production trials. From this "total net N" we subtract the net nitrogen in Sudan hay which, according to the maintenance trials, is 25 per cent of the total nitrogen in Sudan hay. (See p. 535.) This procedure is justified for calculating partial effects because the amount of hay fed for the production trials corresponded approximately to the maintenance level at which the utilization of nitrogen of Sudan hay was measured. We also subtract, from the total net nitrogen, the net nitrogen in molasses beet pulp, assuming that the nitrogen in beet pulp is utilized with the same efficiency as the nitrogen in hay. Admittedly, this is a somewhat wild assumption; but since the amount of nitrogen from beet pulp in the ration is small, even a great relative error in estimating its utilization would not much change the result.

Sample calculation 7 illustrates this calculation. The figures are based on trial 16, cow 1007, body weight 460 kg, metabolic body size 99 kg.

Sample Calculation 7: Partial Net Nitrogen in Casein

1. Total net nitrogen per day:

With a basic nitrogen requirement of 0.172 gram net nitrogen per kg 3 , as explained on p. 534, the cow (1007) in this trial (16) required daily 99×0.172 or 17 grams of net nitrogen. In her milk she secreted daily 43 grams of nitrogen. She stored in her gained body substance 19 grams of nitrogen. Basic nitrogen requirement, added to nitrogen in milk, plus nitrogen in gained body substance, equals total net nitrogen (17 + 43 + 19)....79 grams

2. Partition of net nitrogen in food:

When Sudan hay was fed alone, the total nitrogen was utilized with an efficiency of 25 per cent. (See p. 535.) With this efficiency, the 117 grams total nitrogen in the daily Sudan-hay ration of this trial contributed to the daily net nitrogen 0.25 × 117.....29 grams

3. Partial utilization of nitrogen in casein:

The 47 grams of nitrogen digested (in terms of partial digestibility) from casein fed

per day contained, per 100 grams, $\frac{48}{47} \times 100$ grams of net nitrogen, or.....102 per cent

The partial net nitrogen in *casein* for seven lactation trials of the first series averages 10.5 ± 0.8 grams of net nitrogen per 100 grams of dry casein; the mean efficiency is 77 ± 6 per cent of the total nitrogen.

The partial utilization of the total nitrogen from casein in the second series of trials was 69 and 84 per cent for the two cows. The simple means of all nine

¹⁸ Maynard's (1937, p. 421) formula for calculating the biological value of protein for milk production does not contain the net nitrogen for maintenance; his results should therefore be strictly comparable only within a given plane of nutrition.

trials with casein amount to 10.7 ± 0.6 grams of partial net nitrogen per 100 grams of dry casein and 77 ± 5 grams of partial net nitrogen per 100 grams of total nitrogen in casein. The coefficient of variation in the utilization of nitrogen from casein for these nine trials amounts to \pm 19 per cent.

In their application, these results for the partial nitrogen utilization are possibly limited to our experimental conditions. These conditions, however, were so chosen that they approach reasonably close to practical dairy feeding, at least as far as food level, amount of roughage, and protein ratio are concerned. In this respect our rations may be regarded as reasonably well balanced.

Partial Utilization of Nitrogen from Barley.—The partial net nitrogen in barley was calculated like that in casein (sample calculation 7). From the total net nitrogen (in maintenance, milk, and gain), however, we had to subtract not only the partial net nitrogen from hay (0.25 × total nitrogen in hay) and beet pulp (0.25 × total nitrogen in beet pulp), but also from casein—namely, 77 per cent of the nitrogen furnished in casein according to the partial utilization of nitrogen from casein mentioned in the previous paragraph. The results of two trials of the first series and two trials of the second series averaged 0.96 \pm 0.24 gram of partial net nitrogen per 100 grams of dry barley and 52 \pm 12 grams of partial net nitrogen per 100 grams of total nitrogen in barley.

One trial was carried out with a low-protein ration, feeding barley alone as concentrate in addition to Sudan hay. In this one trial the partial utilization of nitrogen from barley was higher than in any other trial—namely, 81 per cent of the total nitrogen fed in barley. Since the partial digestibility of nitrogen from barley in this trial was 67 per cent, the partial utilization of the partial digested nitrogen was 121 per cent, indicating an increase in the utilization of hay protein by the addition of barley at low-protein level. The higher efficiency in utilizing protein was, however, at the cost of greater waste of energy. Conceivably, therefore, the optimum protein-to-energy ratio in the food may be different for protein utilization and for energy utilization. If this observation should be confirmed, one must conclude that the ideal—that is, the most economical—balance of a ration for animal production is not a biological constant, but depends on the relative costs for protein and food energy.

Partial Utilization of Energy from Glucose.—To derive the partial net energy in glucose, we calculated the total net energy in a trial as the sum of the net energy for maintenance, the energy in milk, and the energy gained as body substance. From this total net energy we subtracted the partial net energy from hay, from beet pulp, and from casein. The remainder is the partial net energy from glucose.

The following calculation illustrates this procedure. The figures are based on trial 16, cow 1007, body weight 460 kg, metabolic body size 99 kg⁸.

Sample Calculation 8: Partial Net Energy in Glucose for Lactation 1. Total net energy per day:

With a basic net-energy requirement per day of 92 kcal per kg³, as explained on p. 537, the cow (1007) in this trial (16) required daily 92 × 99 or 9,108 kcal of net energy for maintenance only. In her milk she gave off daily 6,060 kcal of chemical energy. She now

2. Partition of net energy supplied by feed:

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pulp as for hay, 100 grams of dry matter in molasses beet pulp contains $176 \times \frac{286}{222}$ or 226

3. Partial net energy in glucose:

Since the 2,110 grams of glucose fed daily contained 7,885 kcal total energy, 100 kcal of total energy in glucose supplied $\frac{1,613}{7,885} \times 100$ kcal net energy, or......20.5 per cent

A partial net-energy content in glucose of only 20 per cent of the total energy is the lowest result obtained in our trials. The mean for the first series of trials is 1.41 ± 16 kcal of partial net energy per 100 grams of glucose, or

 38 ± 4 per cent of the total energy. This would amount to $\frac{38}{0.83}$ = 46 per cent

of the partial digested energy in glucose, or $\frac{38}{0.75} = 51$ per cent of the partial

metabolizable energy in glucose. For nine trials of both series the partial net energy in glucose averages 154 ± 15 kcal per 100 grams of glucose, which is 41 ± 4 per cent of the total energy in glucose.

This seems to be a surprisingly low efficiency of energy utilization, particularly of a material like glucose, which needs no change in order to become a normal component of the blood.

The results of twelve trials on steer calves by Mitchell, Hamilton, and Haines (1940) average 49.9 ± 5.4 calories of net energy per 100 calories of metabolizable energy of glucose. The average efficiency of utilization of energy in glucose for lactation in our trials was thus equal to that for growing animals in the trials of Mitchell and his co-workers and does not differ significantly from the efficiency of converting metabolizable energy of starch into energy of body fat in adult steers—namely, 54 ± 2 per cent (Kellner and Köhler, 1900) or 47.8 per cent (Armsby and Fries, 1918). This latter comparison is somewhat surprising. In general the efficiency of energy utilization for milk production appears to be higher than for fattening. In ten trials of Möllgaard (1931, p. 327) with protein ratios (protein net energy divided by net food energy) in a range without effect on energy utilization (from 0.14 to 0.23) the cows produced 1,000 kcal of milk energy from food that in adult steers

would have yielded on the average only 837 kcal of energy in body fat. According to Niels Hansson's (1923, table 4) figures, 100 calories of net energy for fattening in feeds with a low protein content will yield, on the average, 124 calories of net energy in milk production. For high-protein feeds the increased efficiency of total food energy for lactation over that for fattening is still greater because the metabolizability of the digested protein is increased. (See Kleiber, 1929, note 12.) Kriss (1931, p. 160), probably on the basis of the Scandinavian work, concludes that 1 therm of metabolizable energy is equivalent to 0.575 therm of net energy for fattening, but to 0.693 therm of milk energy.

Future investigation should show whether or not the relatively low utilization of the metabolizable energy of glucose (or its high heat increment), observed in our lactation trials, is related to a greater carbon dioxide production in the rumen or to an increased metabolic rate of the cow's own tissues.

Possibly glucose, which can be readily taken into the blood stream, may for this very reason lead to a temporarily excessive metabolic rate. A scheme of such an effect is given by Kleiber (1936). Soskin and Levine (1937) have shown that the rate of "dextrose utilization" increases with increased level of blood sugar. By "utilization" they mean the disappearance of sugar from the blood and extra hepatic tissues "to produce useful energy and heat" (Soskin, 1941). (This terminology is rather unfortunate: not enough is known concerning the usefulness of the form of energy that the animal derives, aside from heat or prior to heat, from the chemical energy of the nutrients.) Soskin thus observed an increased rate of metabolism of sugar with an increased level of blood sugar.

From experiments on man, Carpenter (1940) concluded: "The increases in carbohydrate combustion during the 3 hours following food ingestion were greater, the greater the amounts of reducing and hydrolyzable sugars in the foods, and smaller the greater the amounts of starch or fat in the foods."

It is further to be remembered that we derived the partial efficiency of utilization of energy from glucose by assuming that the partial efficiency of utilization of casein energy was equal to the partial efficiency of utilization of casein nitrogen. This assumption may not be correct. The transfer of food casein to casein in milk involves possibly greater energy losses than nitrogen losses—that is, glucose energy may conceivably be spent in utilizing food casein for lactation.

The partial net energy in the reference substance as a whole, casein plus glucose, was on the average of nine trials (including both series of the experiment) 50 ± 3 per cent of the total, 61 ± 4 per cent of the partial digestible, and 67 ± 4 per cent of the partial metabolizable energy in the reference substance.

Partial Utilization of Energy from Barley.—Sample calculation 9 illustrates the calculation of the partial net energy in barley. The figures are based on trial 18, cow 1007, body weight 473 kg, metabolic body size 101 kg.*.

Sample Calculation 9: Partial Net Energy and Nitrogen-free Net Energy in Barley for Lactation

1. Total net energy per day:

With a basic net-energy requirement per day of 92 kcal per kg\(^{3}\), as explained on p. 537, the cow (1007) in this trial (18) required daily 92×101 or 9.290 kcal of net energy for maintenance only. In her milk she gave off daily 6.360 kcal of chemical energy. She

2. Partition of net energy supplied by feed:

pulp as for hay, 100 grams of dry matter in molasses beet pulp contains $176 \times \frac{286^{19}}{222}$ or 226

keal net energy. The 386 grams of dry molasses beet pulp consumed daily in this trial

Subtracting the net energy supplied by hay, beet pulp, and casein from the total net energy gives the net energy supplied daily by barley: $15,100 - (9,185 + 872 + 1,026) \dots 4,017$ kcal

3. Partial net energy in barley:

The 4,017 keal net energy supplied by 2,396 grams dry barley means $\frac{4,017}{2,396} \times 100$ or 168 kcal net energy per 100 grams dry barley.

The 4,017 kcal net energy supplied by 10,423 kcal total energy in barley means 38 kcal

net energy per 100 kcal total energy in barley.

This means $\frac{3,733}{2,396} \times 100$ or 156 kcal "nitrogen-free" net energy per 100 grams dry barley.

In trial 14 without case in supplement and with 15.4 per cent protein energy in the total food energy, the partial net energy in barley amounted to 156 kcal per 100 grams of dry barley. In four other trials in which the barley was supplemented by case in, so that the protein energy amounted to 18.2 per cent of the total food energy, the partial net energy in barley averaged to 194 ± 25 kcal net energy per 100 grams of dry matter in barley, or to 44 ± 6 per cent of the total, 57 ± 7 per cent of the digestible, and 66 ± 6 per cent of the metabolizable energy in barley.

The calculation of the "nitrogen-free" net energy in barley rests on the assumption that the efficiency of utilizing protein energy is equal to the efficiency of utilizing nitrogen. Sample calculation 9 shows this calculation. The mean result from four trials with casein supplementing the barley amounts to 160 ± 19 kcal "nitrogen-free" net energy per 100 grams of dry barley or 42 ± 5 per cent of the total nitrogen-free food energy in barley.

In the trial in which barley was fed without casein, the nitrogen-free net energy amounted to only 100 kcal per 100 grams of dry barley, or 27 per cent of the total nitrogen-free food energy.

¹⁹ Ratio of metabolizable energy per 100 grams of dry molasses beet pulp and dry hay. ²⁰ According to Niels Hansson, 100 grams of barley contain 170 kcal of net energy for fattening and 211 kcal for milk production.

FOOD VALUE OF BARLEY

The nutritive value of barley with respect to protein and energy can now be calculated as the amount of casein and of glucose that can be replaced by 100 grams of dry matter of barley, provided either of these feeds is a part of a balanced ration for lactation, containing enough Sudan hay to cover the maintenance requirement.

Our experimental data offer three different methods for calculating this casein glucose equivalent of barley.

Calculation from Pair Trials.—The partial net nitrogen per 100 grams of dry barley measured with one cow can be compared with the partial net nitrogen per 100 grams of casein measured in the same trial with the pairmate of the barley cow. This procedure is known as the method of pair trials.

Four quotients
$$\left(\frac{\text{partial net nitrogen in 100 grams barley}}{\text{partial net nitrogen in 100 grams casein}} \times 100\right)$$
 thus ob-

tained average to a mean of 9.4 ± 1.5 grams of casein as equivalent to 100 grams of barley. In the one trial where the barley was not supplemented with casein and where, accordingly, the protein energy ratio in the ration was lower than normal, 100 grams of barley represented a protein equivalent of 15 grams of casein.

The nitrogen-free glucose equivalent of barley measures the nutritive effect of the food energy in barley in addition to the energy already involved in utilizing protein. This nitrogen-free glucose equivalent, calculated for four pair trials (as with the casein equivalent) amounted to 107 ± 18 grams of glucose replaced by the nitrogen-free energy of 100 grams of dry barley. The pair trial with the casein-free barley ration led to a nitrogen-free glucose equivalent of 97 grams of glucose per 100 grams of dry barley.

Calculation from Period Trials.—Instead of comparing the results for two cows in the same period, one may compare the effect of casein in one period with the effect of barley in another period with the same cow. This procedure is called the method of period trials. The partial net nitrogen from barley obtained with cow 1007 in trial 18, for example, is compared with the partial net nitrogen from casein obtained with the same cow in other periods—namely trials 12, 14, 16, and 20. (See table 1.)

Four quotients resulting from this method lead to a mean of 8.9 ± 2.3 grams of casein equivalent per 100 grams of dry barley.

The mean of the four period calculations for the nitrogen-free glucose equivalent amounts to 97 ± 16 grams of nitrogen-free glucose equivalent per 100 grams of dry barley.

Calculation from Combined Pair and Period Trials.—One can use the available data most fully by comparing the partial net nitrogen in barley obtained in our trial with each cow to the partial net nitrogen in casein obtained with both cows in all reference trials for one series. The procedure is as follows:

Sample Calculation 10: Casein-Glucose Equivalent of Barley (Combined Pair and Period Method)

1. Casein equivalent:

Partial net nitrogen per 100 grams dry barley (cow 1021, trial 16)...........0.86 gram

Mean partial net nitrogen per 100 grams dry casein (cow 1021, 3 trials; and cow 1007,

Nitrogen-free glucose equivalent of barley: $\frac{122}{141} \times 100 = 86$ grams per 100 grams dry matter.

3. Casein-glucose equivalent:

100 grams barley dry matter replaces 8.2 grams casein and 86 grams glucose.

TABLE 8
Casein-Glucose Equivalent of Barley

		Replacement equivalent; 100 grams dry barley replace		
Basis of comparison	Protein (casein)	Nitrogen-free energy (glucose)		
	grams	grams		
Chemical composition (total nutrients)	13	100		
Partial digestible nutrients	14	95		
Partial metabolizable nutrients	9	89		
Partial net nutrients:				
from summary, quotients of means	9.0*	104†		
from pair trials, means of quotients	9.4	107		
from period trials, means of quotients	8.9	97		
from combined pair and period trials based on serial means for net nitrogen and net energy.				
means of quotients	8 6	95		

* Calculated as follows:

 $\frac{\text{mean net nitrogen in 100 grams barley}}{\text{mean net nitrogen in 100 grams casein}} \times 100 = \frac{0.96}{10.7} \times 100 = 9.0 \text{ (see p. 539-541)}.$

t Calculated as follows:

 $\frac{\text{mean nitrogen-free net energy in barley}}{\text{mean nitrogen-free net energy in glucose}} \mathbb{I} \times 100 = \frac{160}{154} \times 100 = 104 \text{ (see p. 541-544)},$

The results of the 2-week trials of the first series are based on seven reference trials; those of the 3-week trials of the second series on two reference trials. The simple mean of four quotients amounts to 8.6 ± 2.0 grams of casein equivalent for 100 grams of dry barley. The four nitrogen-free glucose equivalents obtained by the same method of calculation average 95 ± 8 grams glucose equivalent for the nitrogen-free energy in 100 grams of dry barley.

Casein-Glucose Equivalent of Barley.—Table 8 compares the casein-glucose equivalent of barley as calculated on the basis of total, digestible, metabolizable, and net nutrients. The determination of nitrogen content and heat of combustion alone would have led to the same glucose equivalent of barley as the determination of the net energy in our respiration trials. The casein value of barley is higher when based on total or on digestible nitrogen than when

based on net nitrogen; but, in view of the great variability of our results on nitrogen utilization, the difference cannot be regarded as significant.

In our dairy rations 100 grams of dry matter of barley replaced, on the average of both series of respiration trials, 9 grams of cascin and 100 grams of glucose. This casein-glucose equivalent is considerably higher than that obtained in the first series alone and published as preliminary results (Kleiber 1940, table 2).

DISCUSSION

Net Energy vs. Metabolizable Energy.—The nitrogen-free glucose equivalent of barley calculated on the basis of digestible nutrients does not differ from that based on net energy. This result is in line with Kellner's (1919) observation that the "Wertigkeit" of grain was 100 per cent, which means that the utilization of digestible energy from grain for fattening adult steers is equal to the utilization of digestible energy from a pure substance such as starch. To test the significance of feed evaluation according to net energy as compared with metabolizable or digested energy, we should have to measure the partial replacement equivalents or roughages as additions to the maintenance ration, replacing casein and glucose. If we should then find that the casein-glucose equivalent of 1 pound total digestible nutrients in a roughage is significantly lower than that of 1 pound in barley, our finding would justify the evaluation of feeds in terms of net energy rather than digestible energy. This would be true independently of the explanation for the result. Some may attribute to the roughage a higher "work of digestion and absorption," using Zuntz's ideas; or in addition a greater fermentation, as Kellner suggests. Others, abandoning a physiological explanation, may accept Axelsson's (1939) recent merely mathematical concept of a disturbance of the nutritive balance by addition of roughage to the basic food. Regardless of the theory preferred, the fact would remain that 1 pound of total digestible nutrients in roughage added to a given basic ration would produce less milk or body substance than 1 pound in concentrates added to the same basic ration. Our trials thus far have not contributed to the question of "Wertigkeit" of roughage.

The net energy in Sudan hay was 75 per cent of the metabolizable energy.

The net energy in glucose was $\frac{147}{283} \times 100 = 52$ per cent, that of barley

 $\frac{160}{291} \times 100 = 55$ per cent of the metabolizable energy. This fact indicates that

the efficiency of the metabolizable energy in hay for maintenance was greater than the partial efficiency of glucose or barley for lactation.

Paired Respiration Trials vs. Scandinavian Group Trials.—Aside from some results on protein and energy utilization in Sudan hay for maintaining dry cows, and on partial utilization of protein and energy from casein, glucose, and barley for lactation, our trials yield a basis for appraising the relative merits of paired feeding trials with measurement of the nitrogen and carbon balances, as compared with group trials using body weights as criteria for change in body substance.

The coefficient of variation of the net energy in Sudan hay for maintenance is only \pm 8 per cent for both series of our respiration trials.

The partial net energy in glucose, however, is over three times as variable: it has a standard deviation of \pm 29 per cent of the mean. The coefficient of variation for one measurement of the net energy of barley has the same mag-

nitude—namely, ± 24 per cent of the mean.

In thirteen trials carried out by Kellner and Köhler (1900) on adult steers, the standard deviation of the "Ansatzkoeffizienten" of starch (net energy in per cent of metabolizable energy) may be calculated to be \pm 11 per cent of the mean coefficient. This comparison indeed indicates that trials on food utilization for lactation are subjected to greater variability than trials on food utilization by adult steers when equal numbers of animals and equal duration of the experiment are compared. In growing steers, however, according to results of Mitchell, Hamilton, and Haines (1940), the variability of glucose utilization was as great as that with our lactating cows.

Considering the great variability of the results of our respiration trials on lactating cows, one may ask under what conditions an ordinary Scandinavian group trial with dairy cows could yield results as reliable as those obtained

with paired feeding in the respiration chamber.

As discussed before (p. 515), one major error inherent in the ordinary Scandinavian group trial is the estimation of changes in body substance from the changes in body weight. Lush and his co-workers (1928) analyzed the variance in the results of weighing cattle. From the total variance they deducted the partial variance caused by trends in changing weight with time and by differences between individuals; thus they derived a remainder variance that led to a standard error of weight. The mean of this standard error calculated for their data on 238 cows amounts to \pm (5.5 \pm 0.4) kg. This is very close to a standard deviation of \pm 5.9 kg which Kleiber (1929) has calculated from Kellner's and Köhler's (1900) protocols for steers.

With a standard error of \pm 5.5 kg for one weighing, the standard error of the gain in weight, which is the difference between two weights, would be \pm 5.5 $\sqrt{2}$ = \pm 7.8 kg; and the standard error of the mean gain in weight of

six cows would be $\frac{\pm 7.8}{\sqrt{6}}$ = ± 3.2 kg. The difference between the mean gains

in weight of two groups of six cows would therefore be subjected to a standard error of $\pm 3.2\sqrt{2}$ + 4.5 kg. In terms of energy this error would amount to a maximum $4.5\times9,500=\pm43,000$ kcal, assuming the weight error to involve pure fat only.

The milk energy obtained in 10 months from forty-two Jersey and twenty-four Holstein cows at this station (Kleiber and Mead, 1941) averaged 4 million kilocalories per cow. This is 400,000 keal per month. If, then, one ran a Scandinavian group trial with twelve cows (behaving like those discussed above) for only 1 month, the maximum effect of the weight error would amount to only \(\text{11} \) per cent of the result. If the uncertainty of body weight were the only source of error, then the reliability of such a trial of only 1 month's duration would just equal that of one of Kellner's respiration trials with steers and would greatly surpass that of our paired respiration trials with cows. The maximum effect of the weight error of a 6 months' group trial would, according to these figures, amount to only \(\pm 2\) per cent of the result.

This of course applies only to a trial in which the trend in change of weight is kept small; otherwise such a trend would introduce an additional error resulting from the uncertainty of the energy equivalent of change in weight. A lengthy group trial can, however, be so conducted that the trend in weight change is negligible.

Effect of Basic Food on Reliability of Results, and Advantage of Reference Substance.—From four paired trials we can conclude that the "nitrogen-free" net energy for lactation per 100 grams of dry matter is 164 ± 30 kcal for glucose and 160 ± 19 kcal for barley. If the results on glucose were independent of those on barley, the mean difference would be 4 ± 36 .

The mean of the four differences—namely, net energy in barley minus net energy in glucose, each obtained from a pair trial—amounts, however, to 4 ± 18 kcal. The method of pair trials in this case doubled the reliability of the result, as compared with the calculation from single trials, because net energy of barley and glucose within each trial are correlated. One major source of error, whose effect may be greatly reduced by the pair trial, is probably the basic part of the ration, especially the composition and condition of the Sudan hay. The error might have been reduced to one half by use of a less variable maintenance food for the entire investigation—for example, a mixture of Sudan hay and molasses beet pulp ground, prepared in pellet form, and kept in cans. (Compare p. 520.)

Even with an ideal basic food, however, the use of a reference substance would still be preferable to the measurements of absolute values such as net energy, because the effect of other variables may be also reduced by the use of the reference substance. (Compare p. 516.) Until conditions of dairy feeding can be much better defined and standardized than at present, the use of a reference substance for comparisons is a useful tool in measuring food values.

The net energy for lactation in barley was lowest in the low-protein ration. The difference between the result on that ration and the mean of the other results cannot be considered significant, because of the great variability. It is, however, in line with conclusions drawn from earlier work, especially by Möllgaard (1923, 1929); and it tends to support the advantage of using a reference substance with an adjustable protein:energy ratio such as casein and glucose.

The advantage of using these two relatively pure substances as reference substance is obvious even though commercial casein itself is pretty far from having a constant composition. Our trials showed, however, that the advantages of using semipure compounds as reference substance may be outweighed by the effect on the animal's appetite. Palatability seems to be more important in the performance of high-producing dairy cows than in that of fattening steers. Kellner (1900) fattened steers fairly well with rations containing great amounts of pure starch or oil or wheat gluten or even cellulese. We, on the other hand, could not maintain a high lactation in our cows when casein and glucose were substituted for grain and cottonseed meal, even though we trained the cows for months to that diet.

²¹ The correlation coefficient amounts to 0.82; its significance is, however, doubtful, since it is based on only four pairs of results.

Until we can make casein glucose rations nearly as palatable as ordinary feeds, we must conclude that a mixture of a "natural" high-protein and a "natural" low-protein feed, such as cottonseed meal and barley, would probably be a more suitable reference substance for measuring food values with dairy cows than is our mixture of casein and glucose. To obtain barley and cottonseed meal or similar feeds in well-enough-defined composition and condition might be easier than to train high-producing dairy cows sufficiently to casein and glucose.

Cows vs. Steers for Measuring Food Values.—The net energy for lactation in glucose measured in a 3 weeks' respiration trial is about three times as variable as the net energy of starch for fattening measured in a similar trial. This result does not, however, necessitate Möllgaard's apparent conclusion (1927)—namely, that food values for dairy cows should be measured on steers. If the great variability shown in our trials is inherent in food utilization by cows, then the application of any food value to dairy cows would anyway be subjected to an error corresponding to this variability. Applying results from fattening steers to dairy cows would not decrease this random error. Such procedure would, on the contrary, superimpose a systematical error, which originates from the uncertainty of applying the efficiency of food utilization for depositing body fat to a process as different from fattening as is lactation.

For practical feeding trials, dairy cows have a decided advantage. The product of steers in such trials is measured as increase in body weight; and a great source of error, no matter how carefully the trials are conducted, is the interpretation of the increase in weight as increase in energy of body substance. In practical trials with dairy cows, on the contrary, changes in body weight play a minor role. If the trials are carefully conducted, the product (milk) can be measured and analyzed with an accuracy that makes an error from this source negligible compared with other errors of the trial.

There remains a possible source of error inherent in these trials, whether they are conducted with groups in the barn or with pairs in the respiration chamber. Conceivably, cows fed rations higher than their requirement for maintenance and lactation may increase their metabolic rate and thus burn up the extra food taken, instead of using it for gains in body fat and body protein. In this case the added food would seem to have no nutritive value. Whether such a possibility is to be feared more in cows than in steers is not known at present. According to Möllgaard (1923), cows with a reduced milk flow but a continued intake of food at the production level readily gain body substance, and this fattening effect in cows is reliable enough so that "production equivalents" (net energy for fattening per 1,000 kcal of net energy for milk production) can be based on his measurements.

 $^{^{22}}$ The variability of the net energy of glucose measured on growing steers by Mitchell, Hamilton, and Haines (1940) is, however, just as great as the variability of net energy in glucose measured in our lactation trials. The net energy per 100 grams of glucose determined by Mitchell and his co-workers (1940, p. 859) is subject to a coefficient of variation of \pm 39 per cent of the mean. The corresponding coefficient of variation for the net energy of glucose in our lactation trials amounts to \pm 29 per cent of the mean.

CONCLUSIONS

In view of these results, there seems to be no justification for discarding the cows in measuring feed values and particularly no justification for the idea that food values for dairy cows could be measured only with steers.

Through the use of a standardized basic food and a standardized multiple reference substance adjustable to the protein:energy ratio of the feeds to be tested, the Scandinavian group trial should become one of the best methods for measuring food values, especially food values for milk production.

To know for each of the cows used in such group trials the complete carbon and nitrogen balance would be ideal. To run the group trial in twelve respiration chambers, however, not only would be a considerable economic burden, but might involve technical disadvantages because it might introduce artificial conditions. The best improvement over the Scandinavian group trial (with estimation of body gain from body weight) would be the conduction of this trial in two respiration chambers, each housing a group of six cows. That would permit the conduction of the trial under fairly natural conditions and would furnish a reliable basis for measuring the body gains or losses, at least for each group as a whole. This for the measurement of food utilization may be all one needs, whereas for much metabolic research the individual respiration trial remains the only satisfactory procedure.

SUMMARY

In evaluating dairy feeds, one must know how much of one feed replaces a given amount of another as part of a well-balanced dairy ration. This figure is designated as partial replacement equivalent.

The present sources for deriving feed equivalents, such as total digestible nutrients, net energy, starch values, and Scandinavian feed units, lead to very different results and are all open to criticism.

This paper reports an investigation of the possibilities of obtaining better partial replacement equivalents of dairy feeds.

The Scandinavian group trial is at present the most reliable method for measuring replacement equivalents, because the conditions under which these trials are carried out resemble most nearly the conditions under which the results are applied; these trials measure the replacement equivalent of a feed for barley.

The replacement of a single constant reference substance such as barley may lead, however, to a change in the protein:energy ratio, which in turn may affect the efficiency of energy utilization in a dairy ration. The expression of food value in one figure, such as the starch or barley equivalent, moreover, neglects the fact that protein and energy have essentially different physiological functions, and that protein cannot be replaced by nitrogen-free sources of energy.

In paired feeding trials on dairy cows in a double respiration chamber, we measured the daily carbon and nitrogen balances as bases for calculating the changes in body protein and body fat content of the cows. These figures, in addition to protein and energy yield in the milk, are necessary for calculating food utilization.

The daily fasting heat production, measured on two pregnant cows during the fourth and fifth day of fast, amounted to 86 ± 3 kcal per day per kg³⁴, which is about 20 per cent above the level reported for dry empty cows and which, for a 1,000-pound cow, amounts to 8,400 kcal.

The lactation rations consisted of a constant part, fed at the maintenance

level, and a variable or exchangeable part completing the ration.

Sudan hay was used as the constant part of the rations. The nutritive content of this hay for maintenance, together with the maintenance requirement of dry cows, was measured in fourteen 2-week respiration trials (seven trials with two cows each).

The protein in Sudan hay was 62 per cent digestible; and the food energy

was 67 per cent digestible.

An average of 3.5 grams of methane was produced per 100 grams of carbohydrates in the hay, which contained, on the average of seven trials, 222 keal of metabolizable energy per 100 grams of dry matter.

The total nitrogen in Sudan hay was utilized to 25 per cent; the digestible nitrogen to 41 per cent. The energy utilization amounted to 59 per cent of the digestible energy. Dry Sudan hay contained, per 100 grams, 3.3 grams of net protein and 167 kcal of net energy for maintenance.

 Λ 1,000-pound cow would require daily 7 pounds of the Sudan hay to meet her protein requirement, but 14 pounds of the same hay to maintain energy

equilibrium.

In four lactation trials of 3 weeks' duration and five of 2 weeks' duration, a mixture of casein and glucose was added to the Sudan hay to complete a balanced ration containing a well-defined reference substance. In two trials of 3 weeks' duration and three trials of 2 weeks' duration, the glucose and part of the casein were replaced by barley, so that the level of hay remained the same and the protein energy ratio of the entire ration also was unchanged.

Partial effects of feeds added to the Sudan hay are defined as the changes in the results between hay alone and hay with added feeds during lactation. The partial digestibility of nitrogen in casein amounted to only 65 per cent, probably because glucose depresses the digestibility of hay protein. The partial digestibility of energy in glucose averaged 85 per cent. In barley the nitrogen had a partial digestibility of 72 per cent; the energy a partial digestibility of 77 per cent.

The partial methane production of glucose amounted to 1.9 grams of methane per 100 grams of carbohydrates; that of barley, to 3.8 grams.

Casein contained 11 per cent partial net nitrogen; barley 1 per cent.

The partial net energy in glucose averaged 154 kcal per 100 grams of dry matter; that in barley, 194 kcal.

The casein-glucose equivalent of barley has been calculated from our results, arranged as pair trials (comparing simultaneous results on two similar cows), as period trials (comparing the results for each cow at different periods), and as a combination of the two methods (comparing the results of the barley trial with each cow with the serial mean of all results of both cows on the reference substance). On the average, 100 grams of barley will replace 9 grams of casein and 100 grams of glucose. This statement means that the "nitrogen-free" energy in 100 grams of barley is equivalent to 100 grams of glucose.

The coefficient of variation of net energy in Sudan hay for maintenance in each of the two series of experiments amounts to \pm 8 per cent. The results on the partial nutritive effects of easein, glucose, and barley as parts of dairy rations are, however, much more variable. The coefficient of variation for the partial net energy of glucose is \pm 29 per cent; that of barley \pm 24 per cent.

The maximum partial error resulting from the uncertainties of body weight in a well-conducted Scandinavian group trial with twelve cows for 1 month may be estimated as \pm 11 per cent of the mean result. This is less than half the error observed in our paired respiration trials of 2 and 3 weeks' duration. Since the uncertainty of body weight as a criterion for body substance is to be regarded as the major reason for preferring a respiration trial over an ordinary group trial, we must conclude from our results that a well-conducted Scandinavian group trial is at least equal to, and probably superior to, a pair trial in a respiration chamber for measuring replacement equivalents of feeds. Aside from the variability of the results, the group trial has the further advantage of approaching more closely the conditions under which the measured replacement equivalents are to be applied.

The advantage of using semipure materials such as casein and glucose as reference substance may, according to our trials, be overcompensated by the decrease in appetite resulting from these feeds, which was the major difficulty in our experiments, and which led to a loss in body weight as well as an

abnormal decrease in the rate of milk production.

Until the palatability of casein and glucose or a similar semipure reference substance can be considerably increased, a more "natural" substitute reference substance, consisting of a protein-low and protein-high feed, such as a mixture of cottonseed meal and barley, may be preferable as a standard in group trials for measuring replacement equivalents of dairy feeds.

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APPENDIX TABLES

KEY TO APPENDIX TABLE A

Each of the 32 sections of Appendix Table A represents the mean results of ten to fifteen 24-hour respiration trials. The superscript letters in the table constitute a guide to the sequence of the calculations. The meanings of these letters are as follows:

- * Energy in urine = (C in urine) $\times 10$
- ^b C in methane = liters CH₄ × 0.536
- ^e Energy in methane = liters $CH_4 \times 9.5$.
- ^d C in $CO_2 = liters CO_2 \times 0.536$
- $^{\circ}$ C in gained protein = N gained $\times 3.25$
- f Energy in gained protein = N gained × 35.615
- g C in gained fat = total C gained minus C in gained protein
- ^h Energy in gained fat = C in gained fat \times 12.42
- ' Energy in gained body substance = energy in gained fat plus energy in gained protein
- k Heat production = katabolizable energy minus energy in gained body substance

The halves of a pair trial (no. 3) are shown on the next page. In the pages that follow, the halves of the various pair trials are placed side by side.

APPENDIX TABLE A

AVERAGE DAILY NITROGEN, CARBON, AND ENERGY BALANCE*

	Niti	rogen	Carbon		Energy	
	In (gm)	Out (gm)	In (gm)	Out (gm)	In (keal)	Out (kcal)
Trial 3, north chamber; Aug. 7–19, 1	939; cow n	o. 494, wei	ght 467 kg;	$W^{3/4} = 100$	kg ^{3/4}	
Feed: dry Sudan hay, 5,270 gm	103.2		2,222		22,140	
Feces		39.3		693		7,250
Digested nitrogen, carbon, or energy	63.9		1,529		14,890	
Urine		60 0	1	143		1,430
Methane (191 liters)				102b		
	2.0	711.	1 004		11 040	1,820
Metabolizable nitrogen, carbon, or energy	3.9		1,284	*****	11,640	
Milk	1 1212					
Katabolizable nitrogen, carbon, or energy	3.9		1,284		11,640	
Heat (respiration, 2,481 liters CO ₂)				1,330d		$12,230^{1}$
Gain in body substance	3.9			-46		-590 i
Gain in body protein (N gain \times 6.25)	3.9		13 °		140f	
Gain in body fat (C in gained fat \times 1.31)				59 g		-730 [±]
Trial 3, south chamber; Aug. 7–19,	1939; cow	no. 1003, w	veight 493 k	g; W ^{3/4} = 1	05 kg ^{3/4}	
Feed: dry Sudan hay, 5,270 gm	103.2		2,222		22,140	
Feces		34.8		665		6,900
Digested nitrogen, carbon, or energy	68.4		1,557		15.240	
Urine		66.5	1,001	155	10,210	1,550
Methane (184 liters)			1	99b		1.750
Metabolizable nitrogen, carbon, or energy	1.9		1 202		11.040	
			1,303		11,940	
Milk			4 000		** 040	
Katabolizable nitrogen, carbon, or energy	1.9		1,303		11,940	
Heat (respiration, 2,351 liters CO ₂)		****		1,260d		11,410
Gain in body substance	1.9		43		530 ·	
() - i - L - J + - i - / \tag{ \ta} \tag{ \tag} \tag{ \tag{ \tag{ \tag{ \tag{ \tag{ \tag{ \tag{ \tag{ \ta	1 1 0		0.0		mo.4	

1.9

60

70[±]

460h

Gain in body protein (N gain × 6.25)......

Gain in body fat (C in gained fat \times 1.31)....

^{*} Superscript letters are explained in the Key to Appendix Table A, p. 558.

	Nitr	ogen	Carl	bon	Ene	ergy
	In (gm)	Out (gm)	In (gm)	Out (gm)	In (kcal)	Out (kcal)
Trial 5, north chamber; Sept. 4-16	, 1939; cov	v no. 494, w	reight 488 k	$g; W^{3/4} = 1$	04 kg ^{3/4}	
Feed: dry Sudan hay, 4,465 gm. Feces	90.1 58.6 10.1 10.1 10.1	31.5 48.5 	1,855 1,270 1,037 1,037	585 	18,940 12,740 9,700 9,700	6,200 1,400° 1,640° 12,420° -2,720°
Gain in body fat (C in gained fat × 1.31)	****			—248в		-3,080h
Trial 7, north chamber; Oct. 2-13,	1939; cow	no. 1009, w	eight 534 k	g; W ^{3/4} = 1	11kg ^{3/4}	
Feed: dry Sudan hay, 5,715 gm. Feces Digested nitrogen, carbon, or energy Urine. Methane (200 liters). Metabolizable nitrogen, carbon, or energy	115.3 72.0 6.0	43.3	2,433 1,715 1,435	718 173 107 ^b	23,840 16,530 12,900	7,310 1,730a 1,900c
Milk Katabolizable nitrogen, carbon, or energy Heat (respiration, 2,710 liters CO ₂) Gain in body substance Gain in body protein (N gain × 6.25) Gain in body fat (C in gained fat × 1.31)	6.0 6.0 6.0		1,435	1,453 ^d -18	12,900 210 ^f	13,160 ^k -260 ⁱ -470 ^h
Trial 9, north chamber; Oct. 30-Nov.	11, 1939; c	ow no. 1009	, weight 54	0 kg. W ^{3/4} :	= 112 kg ^{3/4}	
Feed: dry Sudan hay, 6,100 gm	123.0 73.0 12.0	50.0 61.0 	2,643 	169 122b	25,770 17,550 13,700 13,700	8,220 1,690a 2,160°
Gain in body substance	12.0 12.0		39 0	-30g	60 i 430 f	-370h

	Niti	ogen	Car	bon	Energy	
	In (gm)	Out (gm)	In (gm)	Out (gm)	In (keal)	Out (kcal)
Trial 5, south chamber; Sept. 4-16,	1939; cow	no. 1003, w	veight 501 k	$g; W^{3/4} = 1$	06 kg ^{3/4}	
Feed: dry Sudan hay, 4,465 gm	90.1		1,853		18,940	1
Feces		31.3		597		5,990
Digested nitrogen, carbon, or energy	58.8		1,258		12,950	
Urine		55.5		139		1,390
Methane (188 liters)				101ъ		1,780
Metabolizable nitrogen, carbon, or energy	3.3		1,018		9,780	
Milk						
Katabolizable nitrogen, carbon, or energy	3.3		1,018		9,780	
Heat (respiration, 2,141 liters CO ₂)				1,148d		11,410
Gain in body substance	3.3			-130		-1,630
Gain in body protein (N gain × 6.25)	3.3	* * * *	110		120f	
Gain in body fat (C in gained fat \times 1.31)				—141 g		$-1,750^{1}$
Feed: dry Sudan hay, 5,312 gm	107.2 69.5	37.7	2,261 1,605	656	22,160 15,360	6,800
	8.6	60 9	1,334	168 103b	11,860	1,680
Methane (192 liters)				103b		1,680 1,820
Methane (192 liters)	8.6		1,334	103b	11,860	1,680 1,820
Methane (192 liters)	8.6 8.6		1,334	103b	11,860	1,680 1,820
Methane (192 liters)	8.6 8.6 8.6		1,334 1,334 67	103b	11,860 11,860 790 ⁱ	1,680 1,820
Methane (192 liters)	8.6 8.6		1,334 1,334 67 28°	103b	11,860 11,860 11,860 790 ¹ 310 ⁴	1,680 1,820 11,070
Milk	8.6 8.6 8.6		1,334 1,334 67	103b	11,860 11,860 790 ⁱ	1,680
Methane (192 liters)	8.6 8.6 8.6 8.6		1,334 1,334 67 28° 39s	103b	11,860 11,860 790 ¹ 310 ⁴ 480 ^h	1,680
Methane (192 liters)	8.6 8.6 8.6 8.6		1,334 1,334 67 28° 39s	103b	11,860 11,860 790 ¹ 310 ⁴ 480 ^h	1,680
Methane (192 liters)	8.6 8.6 8.6 3.6	ow no. 1003	1,334 1,334 67 28° 39s	1,267 ^d	11,860 11,860 	1,680
Methane (192 liters)	8.6 8.6 8.6 8.6 11, 1939; ce	ow no. 1003	1,334 1,334 67 28° 39s 3, weight 51'	103b 1,267d 7 kg; W ^{3/4}	11,860 11,860 	11,070
Methane (192 liters) Metabolizable nitrogen, carbon, or energy Milk Katabolizable nitrogen, carbon, or energy Heat (respiration, 2,364 liters CO2) Gain in body substance. Gain in body protein (N gain × 6.25) Gain in body fat (C in gained fat × 1.31) Trial 9, south chamber; Oct. 30-Nov. Feed: dry Sudan hay, 5,680 gm Feeces Digested nitrogen, carbon, or energy	8.6 8.6 8.6 8.6 11, 1939; ce	ow no. 1003	1,334 1,334 1,334 67 28° 39z 8, weight 51	103b	11,860 	11,070
Methane (192 liters) Metabolizable nitrogen, carbon, or energy Milk Katabolizable nitrogen, carbon, or energy Heat (respiration, 2,364 liters CO ₂) Gain in body substance Gain in body protein (N gain × 6.25) Gain in body fat (C in gained fat × 1.31) Trial 9, south chamber; Oct. 30-Nov. Feed: dry Sudan hay, 5,680 gm Feces. Digested nitrogen, carbon, or energy Jrine.	8.6 8.6 8.6 8.6 11, 1939; c	ow no. 1003	1,334 1,334 1,334 67 28° 392 8, weight 51' 2,461 1,674	103b 1,267d 7 kg; W ^{3/4}	11,860 	11,070
Methane (192 liters) Metabolizable nitrogen, carbon, or energy Milk Katabolizable nitrogen, carbon, or energy Heat (respiration, 2,364 liters CO ₂) Gain in body substance Gain in body protein (N gain × 6.25) Gain in body fat (C in gained fat × 1.31) Trial 9, south chamber; Oct. 30-Nov. Feed: dry Sudan hay, 5,680 gm Feces. Digested nitrogen, carbon, or energy Jrine.	8.6 8.6 8.6 8.6 11, 1939; co	ow no. 1003	1,334 1,334 1,334 67 28° 39s 39s 3, weight 51'	103b 1,267 ^d 7 kg; W ^{3/4} 787	11,860 	1,680 1,820 11,070 11,070 8,040
Methane (192 liters) Metabolizable nitrogen, carbon, or energy Milk Katabolizable nitrogen, carbon, or energy Heat (respiration, 2,364 liters CO ₂) Gain in body substance Gain in body protein (N gain × 6.25) Gain in body fat (C in gained fat × 1.31) Trial 9, south chamber; Oct. 30-Nov. Feed: dry Sudan hay, 5,680 gm Feecs Digested nitrogen, carbon, or energy Jrine Methane (212 liters) Metabolizable nitrogen, carbon, or energy	8.6 8.6 8.6 8.6 11, 1939; co	ow no. 1003	1,334 1,334 1,334 67 28° 39s 3, weight 51 2,461	1,267 ⁴	11,860 	1,680 1,820 11,070 8,040 1,750 2,010
Methane (192 liters)	8.6 8.6 8.6 8.6 11, 1939; co	20w no. 1003	1,334 1,334 1,334 67 28° 39s 3, weight 51 2,461 1,674 1,385	1,267 ⁴	11,860 	1,680 1,820 11,070 8,040 1,750 2,010
Methane (192 liters) Metabolizable nitrogen, carbon, or energy Milk Katabolizable nitrogen, carbon, or energy Heat (respiration, 2,364 liters CO2) Gain in body substance Gain in body protein (N gain × 6.25) Gain in body fat (C in gained fat × 1.31) Trial 9, south chamber; Oct. 30-Nov. Feed: dry Sudan hay, 5,680 gm **eces Digested nitrogen, carbon, or energy Jrine Methane (212 liters) Metabolizable nitrogen, carbon, or energy Milk Katabolizable nitrogen, carbon, or energy	8.6 8.6 8.6 8.6 111, 1939; ce	w no. 1003	1,334 1,334 1,334 67 28° 39° 8, weight 51' 2,461 1,674 1,385	1,2674	11,860 	1,680 1,820 11,070 8,040 1,750 2,010
Methane (192 liters) Metabolizable nitrogen, carbon, or energy Milk Katabolizable nitrogen, carbon, or energy Heat (respiration, 2,364 liters CO2) Gain in body substance Gain in body protein (N gain × 6.25) Gain in body fat (C in gained fat × 1.31) Trial 9, south chamber; Oct. 30-Nov. Feed: dry Sudan hay, 5,680 gm Feeces Digested nitrogen, carbon, or energy Jrine Methane (212 liters) Metabolizable nitrogen, carbon, or energy Milk Katabolizable nitrogen, carbon, or energy	8.6 8.6 8.6 8.6 11, 1939; co	ow no. 1003	1,334 1,334 1,334 1,334 67 28° 39g 8, weight 51' 2,461 1,674 1,385	1,267 ^d	11,860 	1,680 1,820 11,070 11,070 8,040 1,750 2,010
Methane (192 liters) Metabolizable nitrogen, carbon, or energy Milk Katabolizable nitrogen, carbon, or energy Heat (respiration, 2,364 liters CO ₂) Gain in body substance. Gain in body protein (N gain × 6.25) Gain in body fat (C in gained fat × 1.31) Trial 9, south chamber; Oct. 30-Nov. Feed: dry Sudan hay, 5,680 gm Feces. Digested nitrogen, carbon, or energy Urine Methane (212 liters) Metabolizable nitrogen, carbon, or energy Milk Katabolizable nitrogen, carbon, or energy Heat (respiration, 2,628 liters CO ₂)	8.6 8.6 8.6 8.6 11, 1939; co	ow no. 1003	1,334 1,334 1,334 67 28° 39s 3, weight 51 2,461 1,674 1,385	1,267 ⁴	11,860 	1,680 1,820 11,070 8,040 1,750 2,010

	Nitr	ogen	Carb	oon	Ene	rgy
	In (gm)	Out (gm)	In (gm)	Out (gm)	In (kcal)	Out (kcal)
Trial 12, north chamber; Dec. 4-22,	1939; cow	no. 1007, w	veight 463 kg	g; W ^{3/4} = 1	00 kg ^{3/4}	
Dry matter in feed, grams: Sudan hay, 5,296; beet pulp, 530; casein, 529; glucose, 2,109; total dry matter, 8,464	174.0 111.0 67.0 11.0 	63.0 44.0 56.0	3,627 2,640 2,367 1,639 36°	987 	35,060 25,380 21,530 13,790 	9,680 1,280 2,570 8,320 15,880 -2.090
Trial 14, north chamber; Jan. 29-Feb.	10, 1940; c	eow no. 100	7, weight 46	3 kg; W ^{3/4}	= 100 kg ^{3/4}	
Dry matter in feed, grams: Sudan hay, 5,236; beet pulp, 519; casein, 525; glucose, 2,109; total dry matter, 8,389 Feces. Digested nitrogen, carbon, or energy Urine. Methane (298 liters). Methabolizable nitrogen, carbon, or energy Milk, 9.0 kg. Katabolizable nitrogen, carbon, or energy Heat (respiration, 3,335 liters CO ₂) Gain in body substance. Gain in body protein (N gain × 6.25) Gain in body fat (C in gained fat × 1.31)	179.0 114.0 66.0 18.0 	48.0 48.0	3,571 2,585 2,288 1,716 	986 137 160b 572 1,788d -72 	25,390 21,190 14,650	9,660 1,370 2,830 6,540 15,620 -790
Trial 16, north chamber; Feb. 19-Mar	ch 2, 1940	cow no. 10	007, weight 4	160 kg; W ³ /	$^{4} = 99 \text{ kg}^{3/4}$	1
Dry matter in feed, grams: Sudan hay, 5,258; beet pulp, 519; casein, 521; glucose, 2,110; total dry matter, 8,408. Feces. Digested nitrogen, carbon, or energy. Urine. Methane (292 liters). Metabolizable nitrogen, carbon, or energy. Milk, 8.5 kg. Katabolizable nitrogen, carbon, or energy. Heat (respiration, 3,267 liters CO ₂). Gain in body substance. Gain in body protein (N gain × 6.25).	119.0 62.0 19.0	74.0 57.0 43.0	3,534 2,507 2,212 1,683	1,027 138 157b 529 1,751d -68	35, 150 25, 040 20, 890 14, 830	10,110 1,38 2,77 6,06

			1			
	Nitr	ogen	Car	bon	Ene	ergy
	In (gm)	Out (gm)	In (gm)	Out (gm)	In (kcal)	Out (kcal)
Trial 12, south chamber; Dec. 4-22	, 1939; cow	no 1021, w	eight 474 k	g; W ^{3/4} = 1	02 kg ^{3/4}	
Dry matter in feed, grams: Sudan hay, 4,883;						
beet pulp, 530; casein, 529; glucose, 2,109;						
total dry matter, 8,051	166.0		3,448		33,320	
Feces		60.0		944		9,320
Digested nitrogen, carbon, or energy	106.0		2,504		24,000	
Urine		34 0		116		1,160°a
Methane (203 liters)				109Ъ		1,930∘
Metabolizable nitrogen, carbon, or energy	72.0	**···	2,279	*******	20,910	
Milk, 12.5 kg Katabolizable nitrogen, carbon, or energy	20.0	52 0	1 404	795	11 700	9,150
Heat (respiration, 3,288 liters CO ₂)			1,484	1,762d	11,780	15,310k
Gain in body substance	20.0			-278		-3,550°
Gain in body protein (N gain × 6.25)	20.0		650	-210	710 ^f	-0,000
Gain in body fat (C in gained fat \times 1.31)				-343g		-4,260h
	****			0.20-		2,200
Trial 14, south chamber; Jan. 29-Feb.	10, 1940; c	ow no. 1021	l, weight 47	78 kg; W ³ 4	= 102 kg ^{3/4}	
Dry matter in feed, grams: Sudan hay, 5,126;						
beet pulp, 475; barley, 3,281; total dry mat-						
ter, 8,882	164.0		3,791		37,920	
Feces		63 0		1,140		11,280
Digested nitrogen, carbon, or energy	101.0		2,651		26,640	
Urine		46.0		129		1,290*
Methane (329 liters)				176b		3,130∘
Metabolizable nitrogen, carbon, or energy	55.0		2,346		22,220	
Milk, 10.5 kg		48.0		682		7,900
Katabolizable nitrogen, carbon, or energy	7.0		1,664		14,320	
Heat (respiration, 3,409 liters CO ₂)				1,827d		16,380k
Gain in body substance.	7.0			-163	0	-2,060 ⁱ
Gain in body protein (N gain × 6.25)	7.0	,	23 ∘	-186g	250 f	-2,310h
Gain in body fat (C in gained fat \times 1.31)			* * * * * * *	-1908		2,310-
Trial 16, south chamber; Feb. 19-Marc	h 2, 1940; o	cow no. 102	1, weight 4	73 kg; W ^{3/4}	= 101 kg ³ /	4
D						
Dry matter in feed, grams: Sudan hay, 5,258;						
beet pulp, 475; casein, 318; barley, 3,118; total dry matter, 9,169	217.0		3,917		39,540	,,,,,
Feces	217.0	69.0	0,317	1,126		10,980
Digested nitrogen, carbon, or energy	148.0		2,791		28,560	
Urine		74 0		148		1,480a
Methane (345 liters)				185b		3,280°
Metabolizable nitrogen, carbon, or energy	74.0		2,458		2,380	
Milk		49.0		653		7,540
Katabolizable nitrogen, carbon, or energy	25.0		1,805		16,260	
Heat (respiration, 3,411 liters CO ₂)	4.43			1,828ª		16,660k
Gain in body substance	25.0			23		-400i
Gain in body protein (N gain × 6.25)	25.0		810		890f	
Gain in body fat (C in gained fat \times 1.31)				-104s		-1,290b

LEA (C	ontinuea)			
Niti	ogen	Car	bon	Ene	ergy
In (gm)	Out (gm)	In (gm)	Out (gm)	In (kcal)	Out (kcal)
ril 6, 1940;	cow no. 100	7, weight 4	70 kg; W ^{3/4}	$= 101 \text{ kg}^{3/}$	4
175.0 111.0 43.0 	64.0 68.0 44.0 -1.0 -1.0 -1.0 -1.0	3,572 2,490 2,193 1,645	1,082† 136 161b 548 1,689d 443e41g	34,730 	10,530 1,360° 2,860° 6,360 14,170° -550° -40° -510°
163.0 	59.0 44.0 35.0	3,291 2,359 2,097 1,657 32 32	932 	31,990 	9,550 1,250a 2,420c 5,150 13,340k
5, 1940; cov	v no. 1007,	weight 457	kg; W ^{3/4} =	99 kg ^{3/4}	
104.0 64.0 13.0 	40.0 51.0	2,328 1,600 1,364	728 132 10 4 ^b	22,470 15,210 12,050 12,050	7,260 1,320° 1,840°
	Nitr In (gm) 175.0 111.0 43.0 y 5, 1940; co 163.0 25.0 25.0 25.0 5, 1940; cov	Nitrogen In (gm) Out (gm) 175.0 64.0 111.0 68.0 44.01.01.01.01.0 59.0 104.0 44.0 35.0 25.0 35.0 25.0 25.0 25.0 59.0 104.0 44.0 35.0 25.0 59.0	In (gm) Out (gm) In (gm) ril 6, 1940; cow no. 1007, weight 4 175.0 64.0 111.0 2,490 68.0 44.01.0 1,6451.0 1,6451 0 59.0 2,359 44.0 59.0 2,359 44.0 59.0 1,657 25.0 32 25.0 32 25.0 32 25.0 32 25.0 32 25.0 32 25.0 32 25.0 32 25.0 32 25.0 32 25.0 32 25.0 31,657 59.0 32 25.0 32 25.0 32 25.0 32 25.0 31,657 35.0 32 25.0 32 25.0 31,657 35.0 32 25.0 32 25.0 31,657 35.0 32 25.0 32 25.0 32 25.0 31,657 35.0 32 25.0 31,657 35.0 32 25.0 31,657 35.0 32 35.0 32 35.0 32 36.0 32 37.0 35.0 32 38.0 32 38.0 32 39.0 32 30.0 3	Nitrogen Carbon In (gm) Qut (gm) Qut (gm) Out	Nitrogen

 $[\]dagger$ Carbon determination in feces lost, this figure calculated based on equal digestibility of carbon and energy.

			1			
	Niti	ogen	Car	bon	En	ergy
	In (gm)	Out (gm)	In (gm)	Out (gm)	In (keal)	Out (keal)
Trial 18, south chamber; March 25-Apr	ril 6, 1940;	cow no. 102	21, weight 4	173 kg; W ^{3/6}	$s = 101 \text{ kg}^3$	/4
Dry matter in feed, grams: Sudan hay, 5,219; beet pulp, 386; casein, 402; glucose, 1,646; total dry matter, 7,653 Feces Digested nitrogen, carbon, or energy Urine Methane (256 liters) Metabolizable nitrogen, carbon, or energy Milk, 8.0 kg Katabolizable nitrogen, carbon, or energy Heat (respiration, 2,989 liters CO ₂) Gain in body substance Gain in body protein (N gain × 6.25) Gain in body fat (C in gained fat × 1.31) Trial 20, south chamber; April 22-May	99.0 35.0 	38.0 -3.0 -3.0 -3.0	3,275 2,359 2,097 1,587	916‡ 125 137b 510 1,602d -15 -10e -5x	31,350 	9.030 1.250a 2.430c 5.970 12.840k -170c -110f -60h
Dry matter in feed, grams: Sudan hay, 5,274; beet pulp, 395; casein, 402; glucose, 1,646; total dry matter, 7,717. Feces. Digested nitrogen, carbon, or energy. Urine. Methane (252 liters). Metabolizable nitrogen, carbon, or energy. Milk, 7.2 kg. Katabolizable nitrogen, carbon, or energy. Heat (respiration, 2,952 liters CO ₂). Gain in body substance. Gain in body protein (N gain × 6.25). Gain in body fat (C in gained fat × 1.31).	163.0 103.0 54.0 18.0 18.0	60.0 49 0 36.0	3,291 2,337 2,070 1,608	954 	31,990 	9,400 1,320a 2,390c 5,370 13,270k
Trial 22, south chamber; May 13-25	, 1940; cow	no. 1021, v	veight 466 l	(g; W ^{3/4} =	100 kg ^{3/4}	
Feed: dry Sudan hay, 5,318 gm	104.0 62.0 9.0	42.0	2,328 1,570 1,343	758 121 106b	22,470 15,030 11,950	7,440 1,210a 1,870c
$\label{eq:milk} \begin{array}{ll} \mbox{Milk} & \mbox{Katabolizable nitrogen, carbon, or energy} \dots \\ \mbox{Heat (respiration, 2,359 liters CO_2)} & \mbox{Gain in body substance} & \mbox{Gain in body protein (N gain \times 6.25)} & \mbox{Gain in body fat (C in gained fat \times 1.31)} & \mbox{.} \end{array}$	9.0		1,343 79 29° 50g	1,264 ^d	11,950 940 ¹ 320 ^f 620 ^h	11,010k

 $[\]uparrow$ Carbon determination in feces lost, this figure calculated based on 9.86 kcal per gram of carbon in feces of same cow from trials 12 and 20.

	Nitr	rogen	Carl	oon	Ene	rgy
	In (gm)	Out (gm)	In (gm)	Out (gm)	In (keal)	Out (keal)
Trial 24, north chamber; Jan. 27–Feb	. 14, 1941;	cow no. 732	2, weight 40	9 kg; W ^{3/4}	$= 91 \text{ kg}^{3/4}$	
Feed: dry Sudan hay, 4,257 gm	99.3		1,833		18,420	
Peces		36.8		583		6,060
Digested nitrogen, carbon, or energy	62.5		1,250		12,360	
rine		55.2		124		1,240
dethane (149 liters)				80b		1,420
Metabolizable nitrogen, carbon, or energy	7.3		1,046		9,700	
Milk						
Katabolizable nitrogen, carbon, or energy	7.3		1,046		9,700	
Heat (respiration, 2,342 liters CO ₂)				1,255d		12,330
Gain in body substance	7.3			-209		-2,630
Gain in body protein (N gain × 6.25)	7.3		24 0		260f	
Gain in body fat (C in gained fat × 1.31)				-233g		-2,890
Trial 25, north chamber; Feb. 24-Marc	eh 14, 1941	; cow no. 7	32, weight 4	32 kg; W ^{3/4}	$= 95 \text{ kg}^{3/4}$	
Feed: dry Sudan hay, 2,186 gm	50.9	1	936		9,390	
Feces		18.7	1	300		3,240
Digested nitrogen, carbon, or energy	32.2		636		6,150	,,,,,,
Urine		47.0		100		1,000
Methane (111 liters)				60b		1,057
Metabolizable nitrogen, carbon, or energy		-14.8	476		4,090	
Milk		1111			*****	,
Katabolizable nitrogen, carbon, or energy		-14.8	476		4,090	
Heat (respiration, 1,682 liters CO ₂)		14.0		902d		9,310
Gain in body substance		-14.8		-426		-5,220
Gain in body protein (N gain \times 6.25) Gain in body fat (C in gained fat \times 1.31)		-14.8		-48°		52
Gain in body lat (C in gained lat X 1.31)		****		—378g		-4,690
Trial 26, north chamber; March 24-Ap	oril 5, 1 941	; cow no. 7	32, weight 4	34 kg; W ^{3/6}	$4 = 95 \text{ kg}^{3/4}$	
Dry matter in feed, grams: Sudan hay, 2,186; casein, 144; glucose, 1,107; total dry matter,		_				
3,437	72.3		1,455		14,340	
Feces		24 9		341		3,56
Digested nitrogen, carbon, or energy	47.4		1,114	, , , ,	10,780	
Urine		31.0		75		75
Methane (166 liters)				89ъ		1,58
Metabolizable nitrogen, carbon, or energy	16.4		950		8,450	
Milk						
Katabolizable nitrogen, carbon, or energy	16.4		950		8,450	
Heat (respiration, 2,161 liters CO ₂)				1,158d		11,11
Gain in body substance				-208		-2,66
	16.4		53 e		580f	
Gain in body protein (N gain \times 6.25) Gain in body fat (C in gained fat \times 1.31)	10.9		00		000-	

TABI	LE A (C	ontinued	!)			
	Niti	rogen	Car	bon	En	ergy
	In (gm)	Out (gm)	In (gm)	Out (gm)	In (kcal)	Out (keal)
Trial 24, south chamber; Jan 27–Feb	. 14, 1941;	cow no. 72	8, weight 37	2 kg; W ^{3/4}	= 85 kg ^{3/4}	1
Feed: dry Sudan hay, 2,151 gm	50.1		927		9,290	
Feces. Digested nitrogen, carbon, or energy.	21.1	19.0	004	293	0.000	3,000
Urine	31.1	32.9	634	61	6,290	610%
Methane (85 liters)				46h		810
Metabolizable nitrogen, carbon, or energy		-1.8	527		4,870	
Milk						
Katabolizable nitrogen, carbon, or energy		-1.8	527		4,870	
Heat (respiration, 1,620 liters CO ₂)				868d		9,090k
Gain in body substance		-1.8		-341		-4,220i
Gain in body protein (N gain × 6.25)		-1.8		6 e		-60f
Gain in body fat (C in gained fat × 1.31)				-335g		-4,160h
Feed: dry Sudan hay, 3,243 gm. Feces. Digested nitrogen, carbon, or energy. Urine. Methane (125 liters). Metabolizable nitrogen, carbon, or energy. Milk Katabolizable nitrogen, carbon, or energy. Heat (respiration, 1,852 liters CO ₂). Gain in body substance. Gain in body protein (N gain × 6.25). Gain in body fat (C in gained fat × 1.31).	75.5 44.8 6.4 6.4 6.4	30.7	1,398 	86 67 ^b 993 ^d -233	13,980 9,000 6,950 2301	860 A 1,190 c
Trial 26, south chamber; March 24–Apr	ril 15, 1941;	; cow no. 7	28, weight 3	80 kg.; W ³	$4 = 86 \text{ kg}^3$	/4
Dry matter in feed, grams: Sudan hay, 2,186; casein, 144; glucose, 1,107; total dry matter, 3,437. Feces	72.3	31.0	1,455	369	14,340	3,840
Digested nitrogen, carbon, or energy	41.3	04.0	1,086		10,500	600e
Urine		24.6		60 71b		600ª 1,250°
Methane (132 liters)	16.7		955		8,650	
2011	10.7		999		0,000	
Milk	16.7		955		8,650	
Heat (respiration, 2,137 liters CO ₂)	10.4			1,145d		11,090k
Gain in body substance	16.7			- 190		-2,440
Gain in body protein (N gain × 6.25)	16.7		540		590 f	
Gain in body fat (C in gained fat × 1.31)				-244s		-3,030h

	Nitr	ogen	Carb	oon	Energy	
	In (gm)	Out (gm)	In (gm)	Out (gm)	In (keal)	Out (kcal)
Trial 27, north chamber; April 6–18	, 1941; cov	v no. 732, v	veight 430 k	$g; W^{3/4} = 9$	4 kg ^{3/4}	
Dry matter in feed, grams: Sudan hay, 2,186;						
casein, 144; glucose, 1,107; total dry matter,						
3,437	72.3		1,455		14,340	
Feces		28.2		348		3,618
Digested nitrogen, carbon, or energy	44.1		1,107		10,720	
Urine		31 0		77		770
Methane (226 liters)				121 ^b		2,150
Metabolizable nitrogen, carbon, or energy	13.1		909	· · · · · ·	7,800	
Milk	10.1				7 000	
Katabolizable nitrogen, carbon, or energy	13.1		909	1 0114	7,800	11,610
Heat (respiration, 2,259 liters CO ₂)	19 1			1,211 ^d -302		
Gain in body substance	13.1 13.1		43 e		470f	-3,810
Gain in body protein (N gain × 6.25)				-345g		-4,280
Gain in body fat (C in gained fat \times 1.31)		* * * *		-9100		-4,200
Trial 28, south chamber; June 16-Jul	v 2. 1941: (cow no. 728	R. weight 347	kg: W3/4 =	= 80 kg ^{3/4}	
Dry matter in feed, grams: Sudan hay, 3,580;						
beet pulp, 400; casein, 90; barley, 4,183; total dry matter, 8,253	176.8		3,641		35,870	
		69.0	1 1	978		10.090
Feces Digested nitrogen, carbon, or energy	107.8	l .	2,663		25,850	10,020
Urine		28.7		77		770
Methane (262 liters)		20.1		140b		2,490
Metabolizable nitrogen, carbon, or energy	79.1		2,446		22,590	2, 300
Milk, 7.2 kg.		42.5		571		6,410
Katabolizable nitrogen, carbon, or energy	36.6		1,875		16,180	0, 110
Heat (respiration, 2,935 liters CO ₂)	,			1,573d		12,610
Gain in body substance.	36.6		302		3,570i	
Gain in body protein (N gain × 6.25)	36.6		119 e		1,300f	
Gain in body fat (C in gained fat × 1.31)			183g		2,270h	
		1				
Trial 29, south chamber; July 14-Au	g. 1, 1941;	cow no. 72	8, weight 34	7 kg; W ^{3/4}	$= 80 \text{ kg}^{3/4}$	
Dry matter in feed, grams: Sudan hay, 3,580;						
beet pulp, 327; casein, 308; glucose, 2,981;						
total dry matter, 7,196	127.1		3,072		29,490	
Feces		62.2		804		8,110
Digested nitrogen, carbon, or energy	64.9		2,268		21,380	
Urine		27.0		99		990
Methane (260 liters)			,	139ь		2,470
Metabolizable nitrogen, carbon, or energy	37.9		2,030	/	17,920	
Milk, 5.8 kg		36.3		477		5,38
Katabolizable nitrogen, carbon, or energy	1.6		1,553		12,540	
Heat (respiration, 2,930 liters CO ₂)				1,570d		12,75
Gain in body substance	1.6			-17		-21
Gain in body protein (N gain \times 6.25)	1.6		5 e		60 f	
Gain in body fat (C in gained fat \times 1.31)				-22g		-27

TABLE A (Concluded)

Nitrogen	Carbon	Energy
In (gm) Out (gm)	In Out (gm)	In (kcal) Out (kcal)

Trial 27, south chamber; April 6-18, 1941, cow no. 728, weight 382 kg; $W^{3/4} = 86 \text{ kg}^{3/4}$

Dry matter in feed, grams: Sudan hay, 2,186;						
casein, 144; glucose, 1,107; total dry matter,						
3,437	72.3		1,455		14,340	
Feces		28.7		388		4,010
Digested nitrogen, carbon, or energy	43.6		1,067		10,330	
Urine		26.3		65		650a
Methane (185 liters)				99ь		1,760°
Metabolizable nitrogen, carbon, or energy	17.3		903		7,920	
Milk						
Katabolizable nitrogen, carbon, or energy	17.3		903		7,920	
Heat (respiration, 2,236 liters CO ₂)				1,198d		11,660k
Gain in body substance	17.3			-295		-3,740i
Gain in body protein (N gain × 6.25)	17.3		56°		620 f	
Gain in body fat (C in gained fat × 1.31)				-351g		-4,360h

Trial 28, north chamber; June 16–July 2, 1941; cow no. 732, weight 378 kg; $W^{3/4} = 86 \text{ kg}^{3/4}$

Dry matter in feed, grams: Sudan hay, 3,580;						
beet pulp, 400; casein, 376; glucose, 3,232;						
total dry matter, 7,588	138.0		3,231		31,120	
Feces		57.9		794		7,700
Digested nitrogen, carbon, or energy	. 80.1		2,437		23,420	
Urine		27.4		104		1,040a
Methane (248 liters)				133b		2,3600
Metabolizable nitrogen, carbon, or energy	52.7		2,200		20,020	
Milk, 9.1 kg		49.1		757		8,485
Katabolizable nitrogen, carbon, or energy	3.6		1,443		11,540	
Heat (respiration, 2,978 liters CO ₂)				1,596d		13,460k
Gain in body substance	3.6			-153		$-1,920^{\circ}$
Gain in body protein (N gain \times 6.25)	3.6		12 e		128f	
Gain in body fat (C in gained fat × 1.31)				-165g		-2,050h

Trial 29, north chamber; July 14–Aug. 1, 1941; cow no. 732, weight 373 kg; $W^{3/4} = 85 \text{ kg}^{3/4}$

Dry matter in feed, grams: Sudan hay, 3,580; beet pulp, 400; casein, 90; barley, 4,183;						
total dry matter, 8,253	176.8		3,641		35,870	
Feces		66.0		968		9,770
Digested nitrogen, carbon, or energy	110.8		2,673		26,100	
Urine		47.0		135		1,350a
Methane (343 liters)				184b		3,260 ∘
Metabolizable nitrogen, carbon, or energy	63.8		2,354		21,490	
Milk, 8.2 kg		52.1		695		7,850
Katabolizable nitrogen, carbon, or energy	11.7		1,659		13,640	
Heat (respiration, 3,319 liters CO ₂)				1,779d		15,180k
Gain in body substance	11.7			-120		-1,540 i
Gain in body protein (N gain × 6.25)	11.7		38°		420 f	
Gain in body fat (C in gained fat × 1.31)				-158g		-1,960h

APPENDIX TABLE B

SUMMARY OF RESULTS ON FEED UTILIZATION

SUMMARY OF RESULTS ON FEED UTILIZATION
Sudan Hay for Maintenance
a. Protein in Sudan hay
Protein in 100 grams dry hay $(N \times 6.25)$
Digestibility of nitrogen in hay
Digestible protein in 100 grams dry hay
Net nitrogen per 100 grams total nitrogen
Net nitrogen per 100 grams digested nitrogen
Net nitrogen per 100 grams dry Sudan hay
b. Requirement for protein maintenance per day per unit of metabolic body size (W%)
Total nitrogen in Sudan hay
Digested nitrogen in Sudan hay
Dry Sudan hay
Dry Sudan hay for protein maintenance of a 1,000-pound cow per day7.0 pounds
c. Energy in Sudan hay
Heat of combustion in 100 grams Sudan hay423 kcal
Digestibility of energy in Sudan hay
Digestible energy in 100 grams Sudan hay284 ± 2 kcal
Metabolizability of digested energy
Metabolizable energy per 100 grams dry Sudan hay
Net energy in digested energy of Sudan hay
Net energy in metabolizable energy
Net energy in 100 grams dry Sudan hay
d. Requirement for energy maintenance per day per unit of metabolic body size (W%)
Digestible energy
Metabolizable energy
Dry Sudan hay
Dry Sudan hay for energy maintenance of a 1,000-pound dry cow per day13 pounds
e. Methane production from Sudan hay
Methane produced per 100 grams dry Sudan hay
Methane produced per 100 grams dry Sudan hay
Methane produced per 100 grams carbohydrates in Sudan hay 3.5 ± 0.07 grams
Methane produced per 100 grams digestible carbohydrates in Sudan hay $.4.4 \pm 0.08$ grams
1 1 5 6 6 7 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
Barley for Lactation
a. Protein in barley
Protein in 100 grams dry barley
Partial digestibility of nitrogen in barley
Partial digestible nitrogen in 100 grams dry barley
Partial net nitrogen in 100 grams dry barley
b. Energy in barley
Heat of combustion in 100 grams of dry barley
Partial digestibility of energy in barley
Partial digestible energy in 100 grams dry barley
Partial metabolizable energy in 100 grams dry barley
Partial net energy in 100 grams dry barley
c. Methane production from barley
Partial methane production per 100 grams dry barley
Partial methane production per 100 grams carbohydrates in barley3.8 ± 0.6 grams
grams out out out jurates in barrey 3.8 ± 0.0 grams

TABLE B (Concluded)

Casein for Lactation

Total nitrogen in 100 grams dry	casein (as used)
Partial digestibility of nitroge	n in casein
Partial net nitrogen in 100 gra	ms dry casein

Glucose for Lactation

Total energy in 100 grams dry glucose (value for pure glucose)374 kcal
Partial digestibility of energy in glucose
Partial digestible energy in 100 grams dry glucose
Partial methane production from 100 grams glucose
Partial metabolizable energy per 100 grams glucose
Partial net energy per 100 grams glucose

Body weight		Meta- bolic body size									
pounds	kg	kg3/4	pounds	kg	$kg^{3/4}$	pounds	kg	$kg^{3/4}$	pounds	kg	kg3/4
600	272	67	800	363	83	1,000	454	98	1,200	544	113
610	277	68	810	367	84	1,010	458	99	1,210	549	113
620	281	69	820	372	85	1,020	463	100	1,220	553	114
630	286	70	830	376	85	1,030	467	100	1,230	558	115
640	290	70	840	381	86	1,040	472	101	1,240	562	115
650	295	71	850	386	87	1,050	476	102	1,250	567	116
660	299	72	860	390	88	1,060	481	103	1,260	572	117
670	304	73	870	395	88	1,070	485	103	1,270	576	118
680	308	73	880	399	89	1,080	490	104	1,280	581	118
690	313	74	890	404	90	1,090	494	105	1,290	585	119
700	318	75	900	408	91	1,100	499	106	1,300	590	120
710	322	76	910	413	92	1,110	503	106	1,310	594	120
720	327	77	920	417	92	1,120	508	107	1,320	599	121
730	331	78	930	422	93	1,130	513	108	1,330	603	121
740	336	79	940	426	94	1,140	517	108	1,340	608	122
750	340	79	950	431	94	1,150	522	109	1,350	612	123
760	345	80	960	435	95	1,160	526	110	1,360	617	124
770	349	81	970	440	96	1,170	531	111	1,370	621	124
780	354	82	980	445	97	1,180	535	111	1,380	626	125
790	358	82	990	449	97	1,190	540	112	1,390	630	126
800	363	83	1,000	454	98	1,200	544	113	1,400	635	127

